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| (72) Inventor(s)<br>Marlene A Jacobson   | (58) Field of Search<br>UK CL (Edition N ) A5B BHA<br>INT CL <sup>6</sup> A61K 31/52<br>ONLINE: WPI, CLAIMS, DIALOG/BIOTECH   |
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## (54) Inhibition of TNF $\alpha$ production with agonists of the A2b subtype of the adenosine receptor

(57) TNF $\alpha$  production is inhibited by contacting the A2b subtype of the adenosine receptor with an adenosine receptor agonist, especially in monocytes in which cAMP accumulation is increased due to activation of adenylate cyclase. The agonist is preferably adenosine 5'-(N-cyclopropyl)carboxamidoadenosine, 5'-(N-ethyl)carboxamideadenosine, (R)-N<sup>6</sup>-phenyl-2-propyladenosine or cyclohexyladenosine. The agonists may be used in the therapy of autoimmune states. A process for the identification of A2b adenosine receptor agonist, or selective, compounds is described, involving treating monocytes with the compound to determine the degree of TNF $\alpha$  inhibitor, and selecting those compounds which either bind specifically to the A2b adenosine receptor or which include cAMP increase in a cell line expressing the receptor.

GB 2 289 218 A

At least one drawing originally filed was informal and the print reproduced here is taken from a later filed formal copy.

|   |     |
|---|-----|
| 10  | 20  |
| Met Pro Pro Ser Ile Ser Ala Phe Gln Ala Ala Tyr Ile Gly Ile Glu Val Leu Ile Ala | 30  |
|   | 40  |
| Leu Val Ser Val Pro Gly Asn Val Leu Val Ile Trp Ala Val Lys Val Asn Gln Ala Leu | 50  |
|   | 60  |
| Arg Asp Ala Thr Phe Cys Phe Ile Val Ser Leu Ala Val Ala Asp Val Ala Val Gly Ala | 70  |
|   | 80  |
| Leu Val Ile Pro Leu Ala Ile Leu Ile Asn Ile Gly Pro Gln Thr Tyr Phe His Thr Cys | 90  |
|   | 100 |
| Leu Met Val Ala Cys Pro Val Leu Ile Leu Thr Gln Ser Ser Ile Leu Ala Leu Ala     | 110 |
|   | 120 |
| Ile Ala Val Asp Arg Tyr Leu Arg Val Lys Ile Pro Leu Arg Tyr Lys Met Val Val Thr | 130 |
|   | 140 |
| Pro Arg Arg Ala Ala Val Ala Ile Ala Gly Cys Trp Ile Leu Ser Phe Val Val Gly Leu | 150 |
|   | 160 |
| Thr Pro Met Phe Gly Trp Asn Asn Leu Ser Ala Val Glu Arg Ala Trp Ala Ala Asn Gly | 170 |
|   | 180 |
| Ser Met Gly Glu Pro Val Ile Lys Cys Glu Phe Glu Lys Val Ile Ser Met Glu Tyr Met | 190 |
|   | 200 |
| Val Tyr Phe Asn Phe Phe Val Trp Val Leu Pro Pro Leu Leu Leu Met Val Leu Ile Tyr | 210 |
|   | 220 |
| Leu Glu Val Phe Tyr Leu Ile Arg Lys Gln Leu Asn Lys Lys Val Ser Ala Ser Ser Gly | 230 |
|   | 240 |
| Asp Pro Gln Lys Tyr Tyr Gly Lys Glu Leu Lys Ile Ala Lys Ser Leu Ala Leu Ile Leu | 250 |
|   | 260 |
| Phe Leu Phe Ala Leu Ser Trp Leu Pro Leu His Ile Leu Asn Cys Ile Thr Leu Phe Cys | 270 |
|   | 280 |
| Pro Ser Cys His Lys Pro Ser Ile Leu Thr Tyr Ile Ala Ile Phe Leu Thr His Gly Asn | 290 |
|   | 300 |
| Ser Ala Met Asn Pro Ile Val Tyr Ala Phe Arg Ile Gln Lys Phe Arg Val Thr Phe Leu | 310 |
|   | 320 |
| Lys Ile Trp Asn Asp His Phe Arg Cys Gln Pro Ala Pro Pro Ile Asp Glu Asp Leu Pro | 326 |
| Glu Glu Arg Pro Asp Asp   |     |

**F I G. 2A**

atgcggccct 10 ccatctcaggc ttcccaaggcc 30 gcctacatcg gcatcggagg 50 gtcacatcgcc  
ctggtctctg 70 tgccccggaa cgtgctggtg atctgggggg tgaagggtggaa ccaggcccgtg 110  
cggatgcaca 130 ctttctgtctt catcggtgtcg ctggcggtgg ctgtatgtggc cgtgggtggcc 150 170  
ctggtcattcc 190 cccctcgccat cctcatcaac attggccac agaccctactt ccacacctgc  
ctcatggtgtg 250 cctgtccgggt cctcatcctc acccagaggct ccattcctggc cctgctggca 270 290  
attgctgtgg 310 accggctaccc tgggtcaag atccctctcc ggtacaagaat ggtgggtgacc 330 350  
370 ccccgaggcg 390 ctagccggc tgctggatcc tctccttcgt gggtggactg 410  
accctatgt 430 ttggctggaa caatctgagt ggggtggggc gggccctggc agccaacggc 450 470  
490 agcatgggg 510 agcccggtgat caagtgcgg ttcgagaagg tcatcagcat ggagttacatg 530  
550 gtctacttca 570 acttctttgt gtgggtgtcg ccccgcttc tcctcatggt cctcatctac 590  
610 ctggagggtct 630 tctaccataat cgcaaggcag ctaacaaga aggtgtcgcc ctccctccggc 650

2120

670 gaccggcaga agtactatgg gaaggaggctg aagatcgcca agtcgctggc cctcatcctc  
730 ttccttttg ccctcaggctg gctgccttg cacatcctca actgcatac cctcttctgc  
790 ccgtcctgcc acaaagccag catccttacc tacattgcca tcttcctcac gcacggcaac  
850 tcggccatga accccatgt ctatgccttc cgccatccaga agttccgggt cacccttctt  
910 aagatttggaa atgaccattt ccgctgcccag cctgcaccc ccatggacga ggatctccca  
970 gaaaggaggc ctgatgacta g

FIG. 2B

|   |     |
|---|-----|
| Met Pro Ile Met Gly Ser Ser Val Tyr Ile Thr Val Glu Leu Ala Ile Ala Val Leu Ala | 20  |
| 30  | 40  |
| Ile Leu Gly Asn Val Leu Val Cys Trp Ala Val Trp Leu Asn Ser Asn Leu Gln Asn Val | 50  |
| 50  | 60  |
| Thr Asn Tyr Phe Val Val Ser Leu Ala Ala Ala Asp Ile Ala Val Gly Val Leu Ala Ile | 70  |
| 70  | 80  |
| Pro Phe Ala Ile Thr Ile Ser Thr Gly Phe Cys Ala Ala Cys His Gly Cys Leu Phe Ile | 90  |
| 90  | 100 |
| Ala Cys Phe Val Leu Val Leu Thr Gin Ser Ser Ile Phe Ser Leu Leu Ala Ile Ala Ile | 110 |
| 110   | 120 |
| Asp Arg Tyr Ile Ala Ile Arg Ile Pro Leu Arg Tyr Asn Gly Leu Val Thr Gly Thr Arg | 130 |
| 130   | 140 |
| Ala Lys Gly Ile Ile Ala Ile Cys Trp Val Leu Ser Phe Ala Ile Gly Leu Thr Pro Met | 150 |
| 150   | 160 |
| Leu Gly Trp Asn Asn Cys Gly Gln Pro Lys Glu Gly Lys Asn His Ser Gln Gly Cys Gly | 170 |
| 170   | 180 |
| Glu Gly Gln Val Ala Cys Leu Phe Glu Asp Val Val Pro Met Asn Tyr Met Val Tyr Phe | 190 |
| 190   | 200 |
| Asn Phe Phe Ala Cys Val Leu Val Pro Leu Leu Leu Met Leu Gly Val Tyr Leu Arg Ile | 210 |
| 210   | 220 |
| Phe Leu Ala Ala Arg Arg Gln Leu Lys Gln Met Glu Ser Gln Pro Leu Pro Gly Glu Arg | 230 |
| 230   | 240 |
| Ala Arg Ser Thr Leu Gln Lys Glu Val His Ala Ala Lys Ser Leu Ala Ile Ile Val Gly | 250 |
| 250   | 260 |
| Leu Phe Ala Leu Cys Trp Leu Pro Leu His Ile Ile Asn Cys Phe Thr Phe Phe Cys Pro | 270 |
| 270   | 280 |
| Asp Cys Ser His Ala Pro Leu Trp Leu Met Tyr Leu Ala Ile Val Leu Ser His Thr Asn | 290 |
| 290   | 300 |
| Ser Val Val Asn Pro Phe Ile Tyr Ala Tyr Arg Ile Arg Glu Phe Arg Gln Thr Phe Arg | 310 |
| 310   | 320 |
| Lys Ile Ile Arg Ser His Val Leu Arg Gln Gln Glu Pro Phe Lys Ala Ala Gly Thr Ser | 330 |
| 330   | 340 |
| Ala Arg Val Leu Ala Ala His Gly Ser Asp Gly Glu Gln Val Ser Leu Arg Leu Asn Gly | 350 |
| 350   | 360 |
| His Pro Pro Gly Val Trp Ala Asn Gly Ser Ala Pro His Pro Glu Arg Arg Pro Asn Gly | 370 |
| 370   | 380 |
| Tyr Ala Leu Gly Leu Val Ser Gly Gly Ser Ala Gln Glu Ser Gln Gly Asn Thr Gly Leu | 390 |
| 390   | 400 |
| Pro Asp Val Glu Leu Leu Ser His Glu Leu Lys Gly Val Cys Pro Glu Pro Pro Gly Leu | 410 |
| 410   |     |
| Asp Asp Pro Leu Ala Gln Asp Gly Ala Gly Val Ser                                 |     |

5/26

**F I G. 4A**

10 atgccccatca tgggtctc ggtgtacatc acgggtggagc tggccattgc ttgtgtggcc  
70 atcctggcca atgtgttgtt gtgctggcc gtgtgtgtca acagcaacct gcagaacgtc  
130 accaaactact ttgtgtgttc actggggcg gccgacatcg cagtgggtt gctcgccatc  
190 ccctttgcaca tcaccatcag cacccatcgttcc tgcgttgccatc gccacggctg cctcttcatt  
250 gcctgtttcg tcctgttcct cacggcaggcc tccatattca gtctccctggc catcgccatt  
310 gacccgttaca ttggccatccg catcccgctc cggtaacaatg gcttgtgttgc acgtcccatg  
370 gctaaggcca tcattgtccat ctgtgtgggttgc ctgtgttttg ccatcggcct gactccatg  
430 cttagtttga acaaactggg tcaggccaaag gagggaaga accactccca gggctgggg  
490 gaggggccaaat tggcctgtctt ctttgtggat gtgggtccca tgaactacat ggtgtacttc  
550 aacttctttg cctgtgtgtt ggtgtgttgc ctgctcatgc tgggtgttca ttgtcggtatc  
610 ttctgtgggg cggcgttaca gctgtggagc atggagaggcc agcctctggc nnnnnnnnnnnnnnnnn  
650 690 730 770 810 850 890 930 970 1010 1050 1090 1130 1170 1210 1250 1290 1330 1370 1410 1450 1490 1530 1570 1610 1650 1690 1730 1770 1810 1850 1890 1930 1970 2010 2050 2090 2130 2170 2210 2250 2290 2330 2370 2410 2450 2490 2530 2570 2610 2650 2690 2730 2770 2810 2850 2890 2930 2970 3010 3050 3090 3130 3170 3210 3250 3290 3330 3370 3410 3450 3490 3530 3570 3610 3650 3690 3730 3770 3810 3850 3890 3930 3970 4010 4050 4090 4130 4170 4210 4250 4290 4330 4370 4410 4450 4490 4530 4570 4610 4650 4690 4730 4770 4810 4850 4890 4930 4970 5010 5050 5090 5130 5170 5210 5250 5290 5330 5370 5410 5450 5490 5530 5570 5610 5650 5690 5730 5770 5810 5850 5890 5930 5970 6010 6050 6090 6130 6170 6210 6250 6290 6330 6370 6410 6450 6490 6530 6570 6610 6650 6690 6730 6770 6810 6850 6890 6930 6970 7010 7050 7090 7130 7170 7210 7250 7290 7330 7370 7410 7450 7490 7530 7570 7610 7650 7690 7730 7770 7810 7850 7890 7930 7970 8010 8050 8090 8130 8170 8210 8250 8290 8330 8370 8410 8450 8490 8530 8570 8610 8650 8690 8730 8770 8810 8850 8890 8930 8970 9010 9050 9090 9130 9170 9210 9250 9290 9330 9370 9410 9450 9490 9530 9570 9610 9650 9690 9730 9770 9810 9850 9890 9930 9970 10010 10050 10090 10130 10170 10210 10250 10290 10330 10370 10410 10450 10490 10530 10570 10610 10650 10690 10730 10770 10810 10850 10890 10930 10970 11010 11050 11090 11130 11170 11210 11250 11290 11330 11370 11410 11450 11490 11530 11570 11610 11650 11690 11730 11770 11810 11850 11890 11930 11970 12010 12050 12090 12130 12170 12210 12250 12290 12330 12370 12410 12450 12490 12530 12570 12610 12650 12690 12730 12770 12810 12850 12890 12930 12970 13010 13050 13090 13130 13170 13210 13250 13290 13330 13370 13410 13450 13490 13530 13570 13610 13650 13690 13730 13770 13810 13850 13890 13930 13970 14010 14050 14090 14130 14170 14210 14250 14290 14330 14370 14410 14450 14490 14530 14570 14610 14650 14690 14730 14770 14810 14850 14890 14930 14970 15010 15050 15090 15130 15170 15210 15250 15290 15330 15370 15410 15450 15490 15530 15570 15610 15650 15690 15730 15770 15810 15850 15890 15930 15970 16010 16050 16090 16130 16170 16210 16250 16290 16330 16370 16410 16450 16490 16530 16570 16610 16650 16690 16730 16770 16810 16850 16890 16930 16970 17010 17050 17090 17130 17170 17210 17250 17290 17330 17370 17410 17450 17490 17530 17570 17610 17650 17690 17730 17770 17810 17850 17890 17930 17970 18010 18050 18090 18130 18170 18210 18250 18290 18330 18370 18410 18450 18490 18530 18570 18610 18650 18690 18730 18770 18810 18850 18890 18930 18970 19010 19050 19090 19130 19170 19210 19250 19290 19330 19370 19410 19450 19490 19530 19570 19610 19650 19690 19730 19770 19810 19850 19890 19930 19970 20010 20050 20090 20130 20170 20210 20250 20290 20330 20370 20410 20450 20490 20530 20570 20610 20650 20690 20730 20770 20810 20850 20890 20930 20970 21010 21050 21090 21130 21170 21210 21250 21290 21330 21370 21410 21450 21490 21530 21570 21610 21650 21690 21730 21770 21810 21850 21890 21930 21970 22010 22050 22090 22130 22170 22210 22250 22290 22330 22370 22410 22450 22490 22530 22570 22610 22650 22690 22730 22770 22810 22850 22890 22930 22970 23010 23050 23090 23130 23170 23210 23250 23290 23330 23370 23410 23450 23490 23530 23570 23610 23650 23690 23730 23770 23810 23850 23890 23930 23970 24010 24050 24090 24130 24170 24210 24250 24290 24330 24370 24410 24450 24490 24530 24570 24610 24650 24690 24730 24770 24810 24850 24890 24930 24970 25010 25050 25090 25130 25170 25210 25250 25290 25330 25370 25410 25450 25490 25530 25570 25610 25650 25690 25730 25770 25810 25850 25890 25930 25970 26010 26050 26090 26130 26170 26210 26250 26290 26330 26370 26410 26450 26490 26530 26570 26610 26650 26690 26730 26770 26810 26850 26890 26930 26970 27010 27050 27090 27130 27170 27210 27250 27290 27330 27370 27410 27450 27490 27530 27570 27610 27650 27690 27730 27770 27810 27850 27890 27930 27970 28010 28050 28090 28130 28170 28210 28250 28290 28330 28370 28410 28450 28490 28530 28570 28610 28650 28690 28730 28770 28810 28850 28890 28930 28970 29010 29050 29090 29130 29170 29210 29250 29290 29330 29370 29410 29450 29490 29530 29570 29610 29650 29690 29730 29770 29810 29850 29890 29930 29970 30010 30050 30090 30130 30170 30210 30250 30290 30330 30370 30410 30450 30490 30530 30570 30610 30650 30690 30730 30770 30810 30850 30890 30930 30970 31010 31050 31090 31130 31170 31210 31250 31290 31330 31370 31410 31450 31490 31530 31570 31610 31650 31690 31730 31770 31810 31850 31890 31930 31970 32010 32050 32090 32130 32170 32210 32250 32290 32330 32370 32410 32450 32490 32530 32570 32610 32650 32690 32730 32770 32810 32850 32890 32930 32970 33010 33050 33090 33130 33170 33210 33250 33290 33330 33370 33410 33450 33490 33530 33570 33610 33650 33690 33730 33770 33810 33850 33890 33930 33970 34010 34050 34090 34130 34170 34210 34250 34290 34330 34370 34410 34450 34490 34530 34570 34610 34650 34690 34730 34770 34810 34850 34890 34930 34970 35010 35050 35090 35130 35170 35210 35250 35290 35330 35370 35410 35450 35490 35530 35570 35610 35650 35690 35730 35770 35810 35850 35890 35930 35970 36010 36050 36090 36130 36170 36210 36250 36290 36330 36370 36410 36450 36490 36530 36570 36610 36650 36690 36730 36770 36810 36850 36890 36930 36970 37010 37050 37090 37130 37170 37210 37250 37290 37330 37370 37410 37450 37490 37530 37570 37610 37650 37690 37730 37770 37810 37850 37890 37930 37970 38010 38050 38090 38130 38170 38210 38250 38290 38330 38370 38410 38450 38490 38530 38570 38610 38650 38690 38730 38770 38810 38850 38890 38930 38970 39010 39050 39090 39130 39170 39210 39250 39290 39330 39370 39410 39450 39490 39530 39570 39610 39650 39690 39730 39770 39810 39850 39890 39930 39970 40010 40050 40090 40130 40170 40210 40250 40290 40330 40370 40410 40450 40490 40530 40570 40610 40650 40690 40730 40770 40810 40850 40890 40930 40970 41010 41050 41090 41130 41170 41210 41250 41290 41330 41370 41410 41450 41490 41530 41570 41610 41650 41690 41730 41770 41810 41850 41890 41930 41970 42010 42050 42090 42130 42170 42210 42250 42290 42330 42370 42410 42450 42490 42530 42570 42610 42650 42690 42730 42770 42810 42850 42890 42930 42970 43010 43050 43090 43130 43170 43210 43250 43290 43330 43370 43410 43450 43490 43530 43570 43610 43650 43690 43730 43770 43810 43850 43890 43930 43970 44010 44050 44090 44130 44170 44210 44250 44290 44330 44370 44410 44450 44490 44530 44570 44610 44650 44690 44730 44770 44810 44850 44890 44930 44970 45010 45050 45090 45130 45170 45210 45250 45290 45330 45370 45410 45450 45490 45530 45570 45610 45650 45690 45730 45770 45810 45850 45890 45930 45970 46010 46050 46090 46130 46170 46210 46250 46290 46330 46370 46410 46450 46490 46530 46570 46610 46650 46690 46730 46770 46810 46850 46890 46930 46970 47010 47050 47090 47130 47170 47210 47250 47290 47330 47370 47410 47450 47490 47530 47570 47610 47650 47690 47730 47770 47810 47850 47890 47930 47970 48010 48050 48090 48130 48170 48210 48250 48290 48330 48370 48410 48450 48490 48530 48570 48610 48650 48690 48730 48770 48810 48850 48890 48930 48970 49010 49050 49090 49130 49170 49210 49250 49290 49330 49370 49410 49450 49490 49530 49570 49610 49650 49690 49730 49770 49810 49850 49890 49930 49970 50010 50050 50090 50130 50170 50210 50250 50290 50330 50370 50410 50450 50490 50530 50570 50610 50650 50690 50730 50770 50810 50850 50890 50930 50970 51010 51050 51090 51130 51170 51210 51250 51290 51330 51370 51410 51450 51490 51530 51570 51610 51650 51690 51730 51770 51810 51850 51890 51930 51970 52010 52050 52090 52130 52170 52210 52250 52290 52330 52370 52410 52450 52490 52530 52570 52610 52650 52690 52730 52770 52810 52850 52890 52930 52970 53010 53050 53090 53130 53170 53210 53250 53290 53330 53370 53410 53450 53490 53530 53570 53610 53650 53690 53730 53770 53810 53850 53890 53930 53970 54010 54050 54090 54130 54170 54210 54250 54290 54330 54370 54410 54450 54490 54530 54570 54610 54650 54690 54730 54770 54810 54850 54890 54930 54970 55010 55050 55090 55130 55170 55210 55250 55290 55330 55370 55410 55450 55490 55530 55570 55610 55650 55690 55730 55770 55810 55850 55890 55930 55970 56010 56050 56090 56130 56170 56210 56250 56290 56330 56370 56410 56450 56490 56530 56570 56610 56650 56690 56730 56770 56810 56850 56890 56930 56970 57010 57050 57090 57130 57170 57210 57250 57290 57330 57370 57410 57450 57490 57530 57570 57610 57650 57690 57730 57770 57810 57850 57890 57930 57970 58010 58050 58090 58130 58170 58210 58250 58290 58330 58370 58410 58450 58490 58530 58570 58610 58650 58690 58730 58770 58810 58850 58890 58930 58970 59010 59050 59090 59130 59170 59210 59250 59290 59330 59370 59410 59450 59490 59530 59570 59610 59650 59690 59730 59770 59810 59850 59890 59930 59970 60010 60050 60090 60130 60170 60210 60250 60290 60330 60370 60410 60450 60490 60530 60570 60610 60650 60690 60730 60770 60810 60850 60890 60930 60970 61010 61050 61090 61130 61170 61210 61250 61290 61330 61370 61410 61450 61490 61530 61570 61610 61650 61690 61730 61770 61810 61850 61890 61930 61970 62010 62050 62090 62130 62170 62210 62250 62290 62330 62370 62410 62450 62490 62530 62570 62610 62650 62690 62730 62770 62810 62850 62890 62930 62970 63010 63050 63090 63130 63170 63210 63250 63290 63330 63370 63410 63450 63490 63530 63570 63610 63650 63690 63730 63770 63810 63850 63890 63930 63970 64010 64050 64090 64130 64170 64210 64250 64290 64330 64370 64410 64450 64490 64530 64570 64610 64650 64690 64730 64770 64810 64850 64890 64930 64970 65010 65050 65090 65130 65170 65210 65250 65290 65330 65370 65410 65450 65490 65530 65570 65610 65650 65690 65730 65770 65810 65850 65890 65930 65970 66010 66050 66090 66130 66170 66210 66250 66290 66330 66370 66410 66450 66490 66530 66570 66610 66650 66690 66730 66770 66810 66850 66890 66930 66970 67010 67050 67090 67130 67170 67210 67250 67290 67330 67370 67410 67450 67490 67530 67570 67610 67650 67690 67730 67770 67810 67850 67890 67930 67970 68010 68050 68090 68130 68170 68210 68250 68290 68330 68370 68410 68450 68490 68530 68570 68610 68650 68690 68730 68770 68810 68850 68890 68930 68970 69010 69050 69090 69130 69170 69210 69250 69290 69330 69370 69410 69450 69490 69530 69570 69610 69650 69690 69730 69770 69810 69850 69890 69930 69970 70010 70050 70090 70130 70170 70210 70250 70290 70330 70370 70410 70450 70490 70530 70570 70610 70650 70690 70730 70770 70810 70850 70890 70930 70970 71010 71050 71090 71130 71170 71210 71250 71290 71330 71370 71410 71450 71490 71530 71570 71610 71650 71690 71730 71770 71810 71850 71890 71930 71970 72010 72050 72090 72130 72170 72210 72250 72290 72330 72370 72410 72450 72490 72530 72570 72610 72650 72690 72730 72770 72810 72850 72890 72930 72970 73010 73050 73090 73130 73170 73210 73250 73290 73330 73370 73410 73450 73490 73530 73570 73610 73650 73690 73730 73770 73810 73850 73890 73930 73970 74010 74050 74090 74130 74170 74210 74250 74290 74330 74370 74410 74450 74490 74530 74570 74610 74650 74690 74730 74770 74810 74850 74890 7493

**FIG. 4B**

670 gcacggtc  
690 cactgcagaa ggagggtccat gctgccaagt cactggccat catttgtggg  
710 730 750 770  
ctctttggcc tctgttggtt gcccctaac atcatcaact gcttcaactt ctctgtcccc  
790 810 830  
gactgcaggc acgccccctt ctggctcatg tacctggcca tcgttccctctc ccacaccaat  
850 870 890  
tcgttgttga atcccttcat ctacgcctac cgttatccgg agttccggca gacccttcggc  
910 930 950  
aagatcattc gcagccacgt cctgaggcag caagaacctt tcaaggcagg tggcaccagt  
970 990 1010  
gccccgggttct tggcagctca tggcagtgtac ggagaggcagg tcagccttcg tctcaacggc  
1030 1050 1070  
cacccggccag gagtgtggc caacggcagt gctcccccacc ctgaggcgag gcccaatggc  
1090 1110 1130  
tatgccctgg ggctgttgtag tggaggaggat gcccaaggat cccaggggaa cacggggctc  
1150 1170 1190  
ccagacgtgg agctcccttag ccatgaggctc aaggaggatgt gcccaggaggcc ccctggcccta  
1210 1230  
gatgaccccc tggcccaaggta tggaggcagga tggtccctga

1/20

Met Leu Leu Glu Thr Gln Asp Ala Leu Tyr Val Ala Leu Glu Leu Val Ile Ala Ala Leu  
 Ser Val Ala Gly Asn Val Leu Val Cys Ala Ala Val Gly Thr Ala Asn Thr Leu Gln Thr  
 Pro Thr Asn Tyr Phe Leu Val Ser Leu Ala Ala Asp Val Ala Val Gly Leu Phe Ala  
 Ile Pro Phe Ala Ile Thr Ile Ser Leu Gly Phe Cys Thr Asp Phe Tyr Gly Cys Leu Phe  
 Leu Ala Cys Phe Val Leu Val Leu Thr Gln Ser Ser Ile Phe Ser Leu Leu Ala Val Ala  
 Val Asp Arg Tyr Leu Ala Ile Cys Val Pro Leu Arg Tyr Lys Ser Leu Val Thr Gly Thr  
 Arg Ala Arg Gly Val Ile Ala Val Leu Trp Val Leu Ala Phe Gly Ile Gly Leu Thr Pro  
 Phe Leu Gly Trp Asn Ser Lys Asp Ser Ala Thr Asn Asn Cys Thr Glu Pro Trp Asp Gly  
 Thr Thr Asn Glu Ser Cys Cys Leu Val Lys Cys Leu Phe Glu Asn Val Val Pro Met Ser  
 Tyr Met Val Tyr Phe Asn Phe Phe Gly Cys Val Leu Pro Pro Leu Leu Ile Met Leu Val  
 Ile Tyr Ile Lys Ile Phe Leu Val Ala Cys Arg Gln Leu Gln Arg The Glu Leu Met Asp  
 His Ser Arg Thr Thr Leu Gln Arg Glu Ile His Ala Ala Lys Ser Leu Ala Met Ile Val  
 Gly Ile Phe Ala Leu Cys Trp Leu Pro Val His Ala Val Asn Cys Val Thr Leu Phe Gln  
 Pro Ala Gln Gly Lys Asn Lys Pro Lys Trp Ala Met Asn Met Ala Ile Leu Leu Ser His  
 Ala Asn Ser Val Val Asn Pro Ile Val Tyr Ala Tyr Arg Asn Arg Asp Phe Arg Tyr Thr  
 Phe His Lys Ile Ile Ser Arg Tyr Leu Leu Cys Gln Ala Asp Val Lys Ser Gly Asn Gly  
 Gln Ala Gly Val Gln Pro Ala Leu Gly Val Gly Leu

FIG. 5

8/20

|     |             |             |             |              |              |             |
|-----|-------------|-------------|-------------|--------------|--------------|-------------|
| 10  | atgtgttgtt  | agacacaggaa | cggcgctgtac | gtggcgctgg   | aggctgggttt  | cgcccgccgtt |
| 70  | tgcggggggg  | gcaaacgtgtt | gttgtgtcgcc | gggggtggca   | cgggcaaacac  | tctgcagacg  |
| 130 | ccccacaact  | acttccttgtt | gtcccctggct | ggggcccgacg  | tggccgtggg   | gctcttcgcc  |
| 190 | atccctttg   | ccatcaccat  | cagcctggcc  | ttctgcacttg  | acttctacgg   | ctgcctcttc  |
| 250 | ctcgccctgtt | tcgtgttgtt  | gctcacgcag  | agtcacatct   | ttagccatct   | ggccgtggca  |
| 310 | gtcgacagat  | acctggccat  | ctgtgtcccg  | ctcaggataa   | aaagttttgtt  | cacggggacc  |
| 370 | cgagcaagag  | gggtcattgtc | tgtccctctgg | gtcccttgccct | ttggcatcggtt | attgactcca  |
| 430 | ttcctgggtt  | ggaacagtaa  | agacagtgcc  | accaacaact   | gcacagaacc   | ctgggatggaa |
| 490 | accacgaatg  | aaagctgtctg | ccttgtaag   | tgtctcttttg  | agaatgttgtt  | ccccatggcc  |
| 550 | tacatggat   | atttcaattt  | ctttgggtgt  | gttctggccct  | cactgcttat   | aatgctgggtt |
| 610 | atctacatta  | agatcttcctt | gtggcctgc   | aggcaggcttc  | aggcggactgaa | gctgtatggac |

FIG. 6A

9/20

670 cactcgagga ccacccctcca gccccggatc catgcaggcca agtcaactggc catgatttgt  
730 gggattttg ccctgtgtcg gttacctgtg catgtgttta actgtgtcac tcttttccag  
790 ccagctcagg gtaaaaaataa gcccaagggg gcaatgaata tggccattt tctgtcacat  
850 gccaaatttag ttgtcaatcc cattgtctat gcttaccggg accggagactt ccgctacact  
910 tttcacaaaa ttatctccag gtatcttc tgccaaagg atgtcaagag tggaaatggt  
970 caggctgggg tacagccctgc tctcggttgt ggcctatga

FIG. 6B

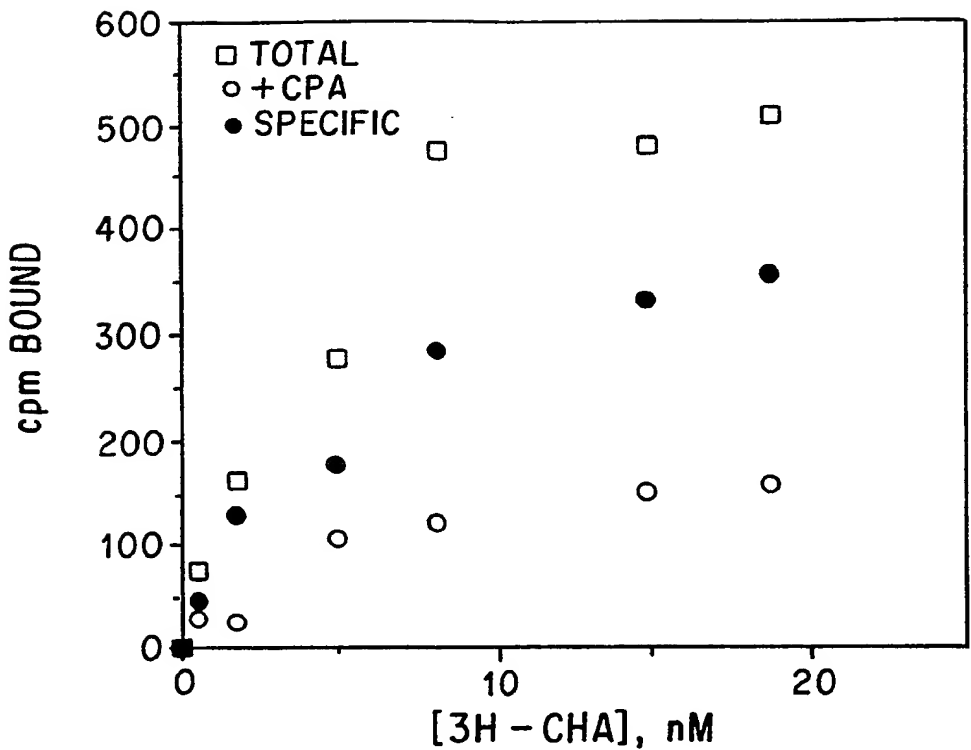


FIG. 7

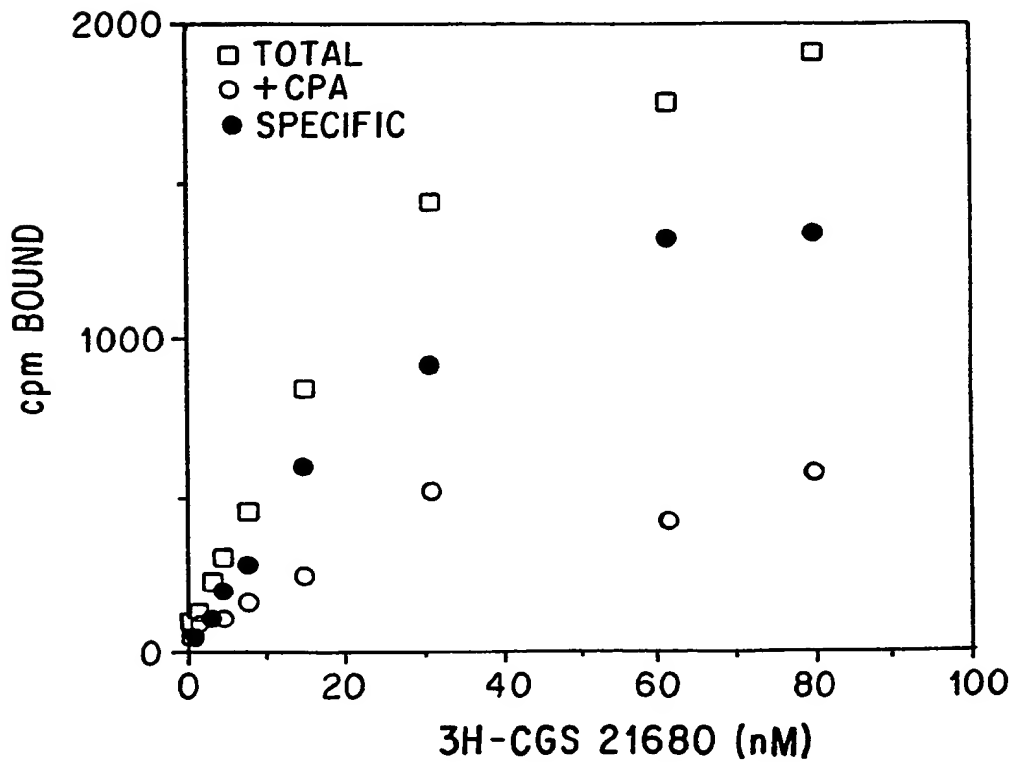


FIG. 8

|   |     |     |
|---|-----|-----|
|   | 10  | 20  |
| Met Pro Asn Asn Ser Thr Ala Leu Ser Leu Ala Asn Val Thr Tyr Ile Thr Met Glu Ile |     |     |
|   | 30  | 40  |
| Phe Ile Gly Leu Cys Ala Ile Val Gly Asn Val Leu Val Ile Cys Val Val Lys Leu Asn |     |     |
|   | 50  | 60  |
| Pro Ser Leu Gln Thr Thr Phe Tyr Ile Val Ser Leu Ala Leu Ala Asp Ile Ala         |     |     |
|   | 70  | 80  |
| Val Gly Val Leu Val Met Pro Leu Ala Ile Val Val Ser Leu Gly Ile Thr Ile His Phe |     |     |
|   | 90  | 100 |
| Tyr Ser Cys Leu Phe Met Thr Cys Leu Leu Leu Ile Phe Thr His Ala Ser Ile Met Ser |     |     |
|   | 110 | 120 |
| Leu Leu Ala Ile Ala Val Asp Arg Tyr Leu Arg Val Lys Leu Thr Val Arg Tyr Lys Arg |     |     |
|   | 130 | 140 |
| Val Thr Thr His Arg Arg Ile Trp Leu Ala Leu Gly Leu Cys Trp Leu Val Ser Phe Leu |     |     |
|   | 150 | 160 |
| Val Gly Leu Thr Pro Met Phe Gly Trp Asn Met Lys Leu Thr Ser Glu Tyr His Arg Asn |     |     |
|   | 170 | 180 |
| Val Thr Phe Leu Ser Cys Gln Phe Val Ser Val Met Arg Met Asp Tyr Met Val Tyr Phe |     |     |
|   | 190 | 200 |
| Ser Phe Leu Thr Trp Ile Phe Ile Pro Leu Val Val Met Cys Ala Ile Tyr Leu Asp Ile |     |     |
|   | 210 | 220 |
| Phe Tyr Ile Ile Arg Asn Lys Leu Ser Leu Asn Leu Ser Asn Ser Lys Glu Thr Gly Ala |     |     |
|   | 230 | 240 |
| Phe Tyr Gly Arg Glu Phe Lys Thr Ala Lys Ser Leu Phe Leu Val Leu Phe Leu Ala     |     |     |
|   | 250 | 260 |
| Leu Ser Trp Leu Pro Leu Ser Ile Ile Asn Cys Ile Ile Tyr Phe Asn Gly Glu Val Pro |     |     |
|   | 270 | 280 |
| Gln Leu Val Leu Tyr Met Gly Ile Leu Leu Ser His Ala Asn Ser Met Met Asn Pro Ile |     |     |
|   | 290 | 300 |
| Val Tyr Ala Tyr Lys Ile Lys Lys Phe Lys Glu Thr Tyr Leu Leu Ile Leu Lys Ala Cys |     |     |
|   | 310 |     |
| Val Val Cys His Pro Ser Asp Ser Leu Asp Thr Ser Ile Glu Lys Asn Ser Glu         |     |     |

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# FIG. 10A

10 atgccccaaa 30 gccaaatgttgcctacatcac 50 catggaaatt  
70 ttcattggac 90 gtggggcaac 110 caagctgaac  
130 cccaggctgc 150 cttcattttc 170 tgacattgtc  
190 gttgggtgtgc 210 ttggccatt 230 aatccacttc  
250 tacagctgcc 270 ttgcctactg 290 catcatgtcc  
310 ttcgtggcca 330 cgggtcaaggc 350 atacaaggagg  
370 gtcaccactc 390 ttaccgttag 410 gtcattccctg  
430 acagaagaat 450 atggctggcc 470 gctggcttgt  
490 gttggattga 510 tggctggaaac 530 ccacagaat  
550 gtcaccttcc 570 atttgcattcc 590 ggtatacttc  
610 agcttcctca 630 cttggatttt 650 tcttgacatc  
tttacatca 650 ttcgaaacaa 650 actcagtctg aacttatcta actccaaaga gacagggtgca

|     |            |             |             |             |              |             |
|-----|------------|-------------|-------------|-------------|--------------|-------------|
| 670 | tttatggac  | gggaggttcaa | gacggcttaag | tccttgc     | tggttttc     | tgtttttgt   |
| 730 | cgtcatggc  | tgcctttatc  | tatcatcaac  | tgcattatct  | actttaatgg   | tgaggtacca  |
| 790 | cagcttgtgc | tgtacatgg   | catccatgtcg | tcccattgc   | actccatgtatc | gaaccctatc  |
| 850 | gtctatgcct | ataaaataaa  | gaagttcaag  | gaaacctacc  | ttttgatcct   | caaaggcctgt |
| 910 | gtggtctgcc | atccctctga  | ttctttggac  | acaaggcattg | agaagaattc   | tgagttag    |

## FIG. 10B

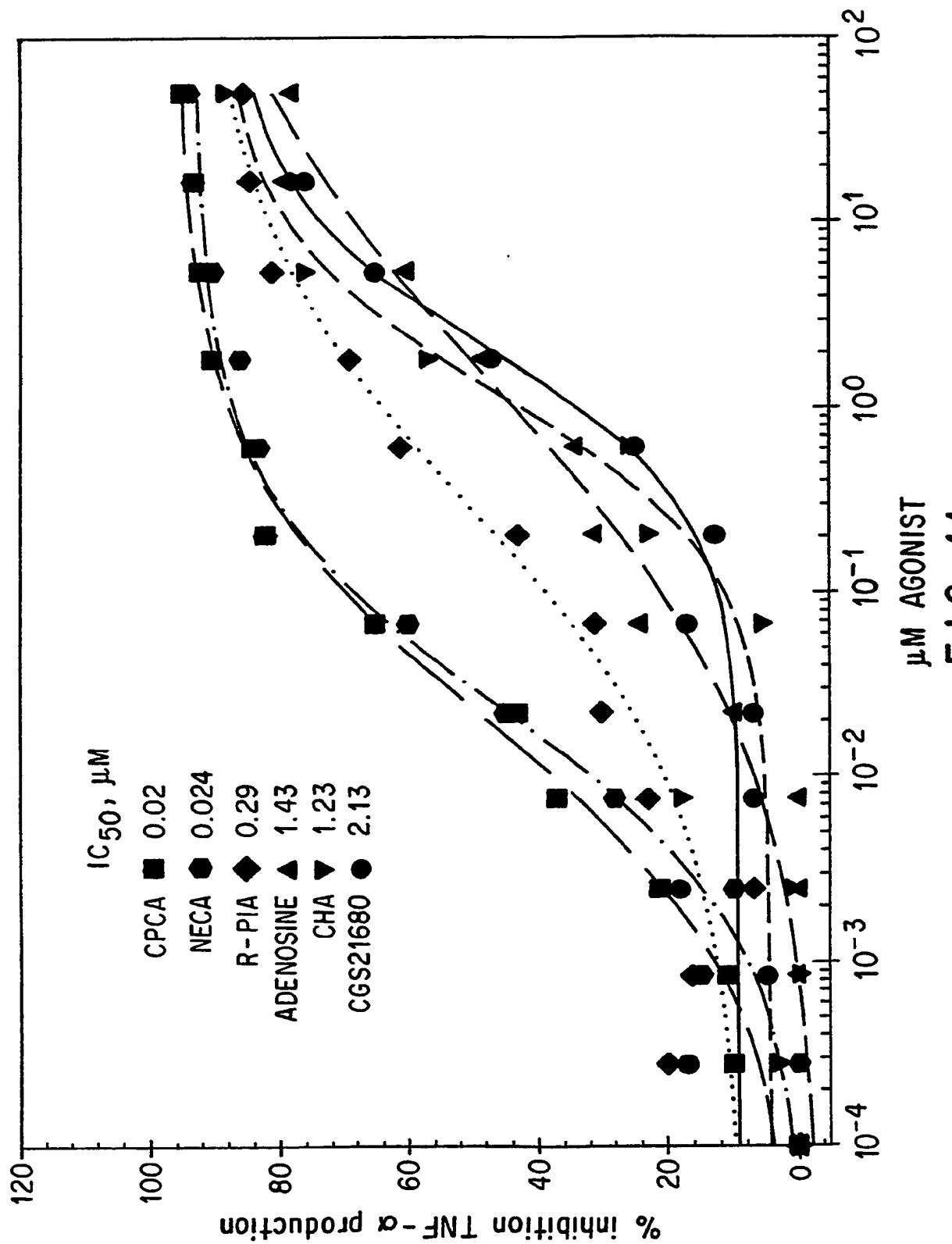
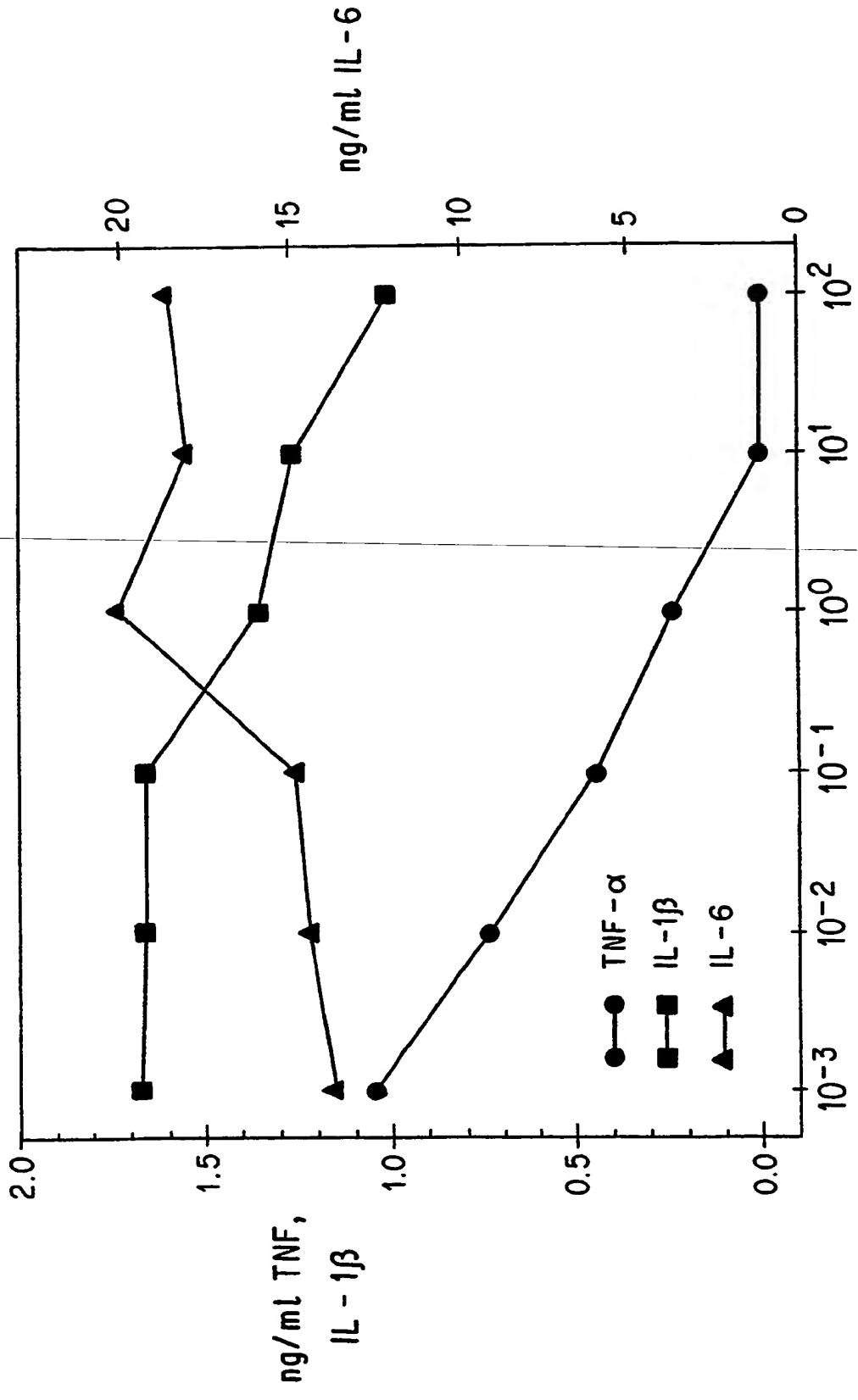


FIG. 11

FIG. 12



16 / 20

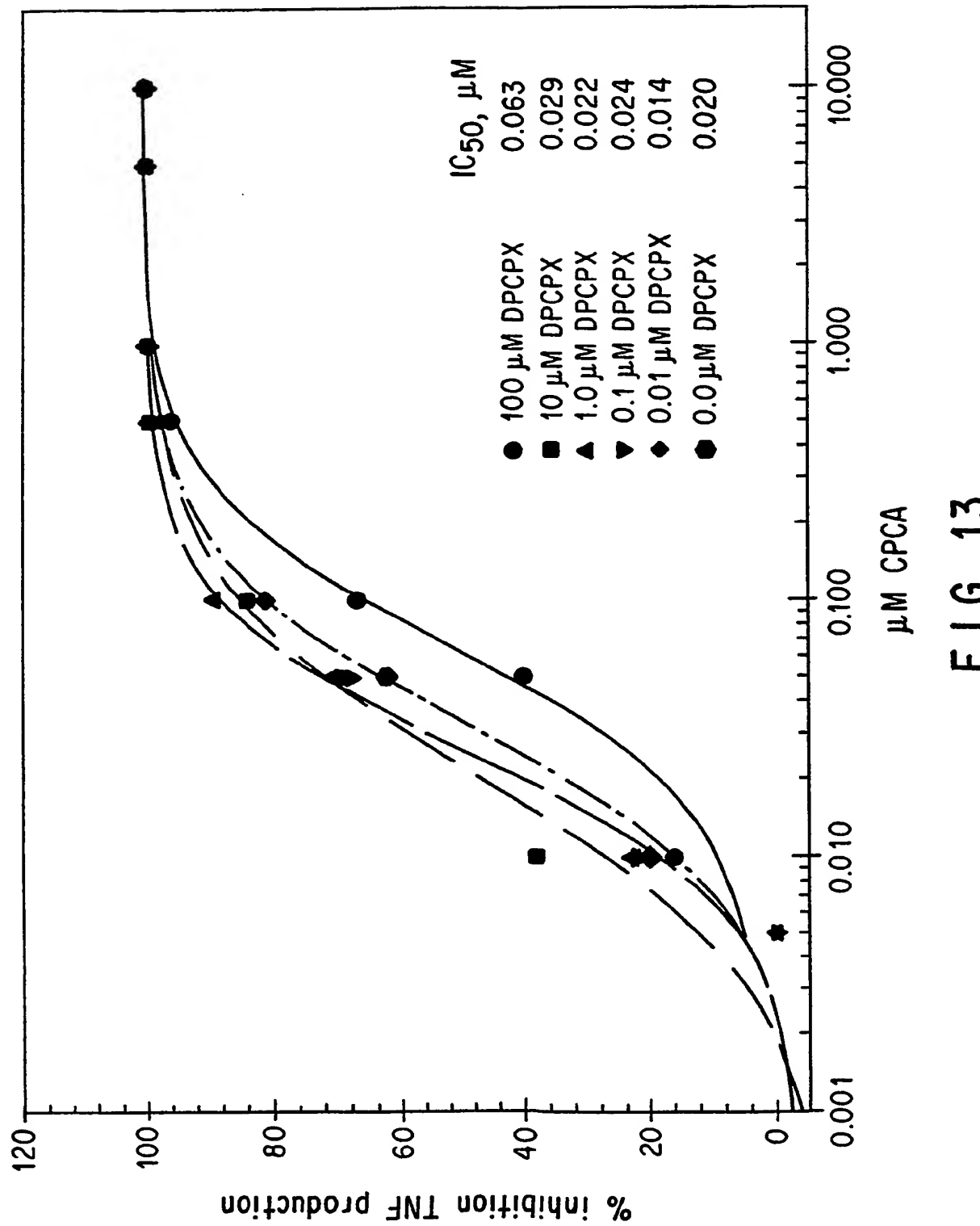
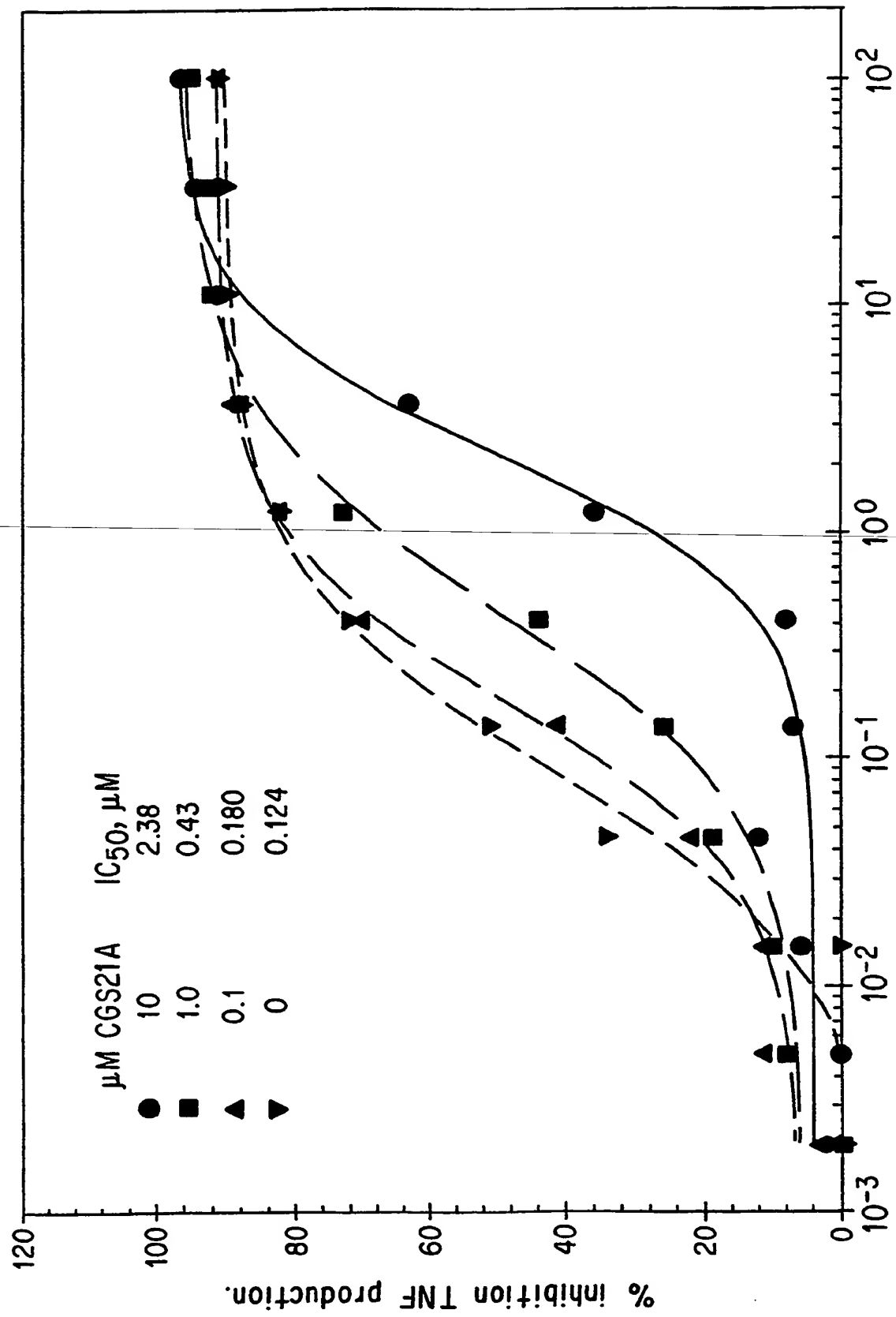
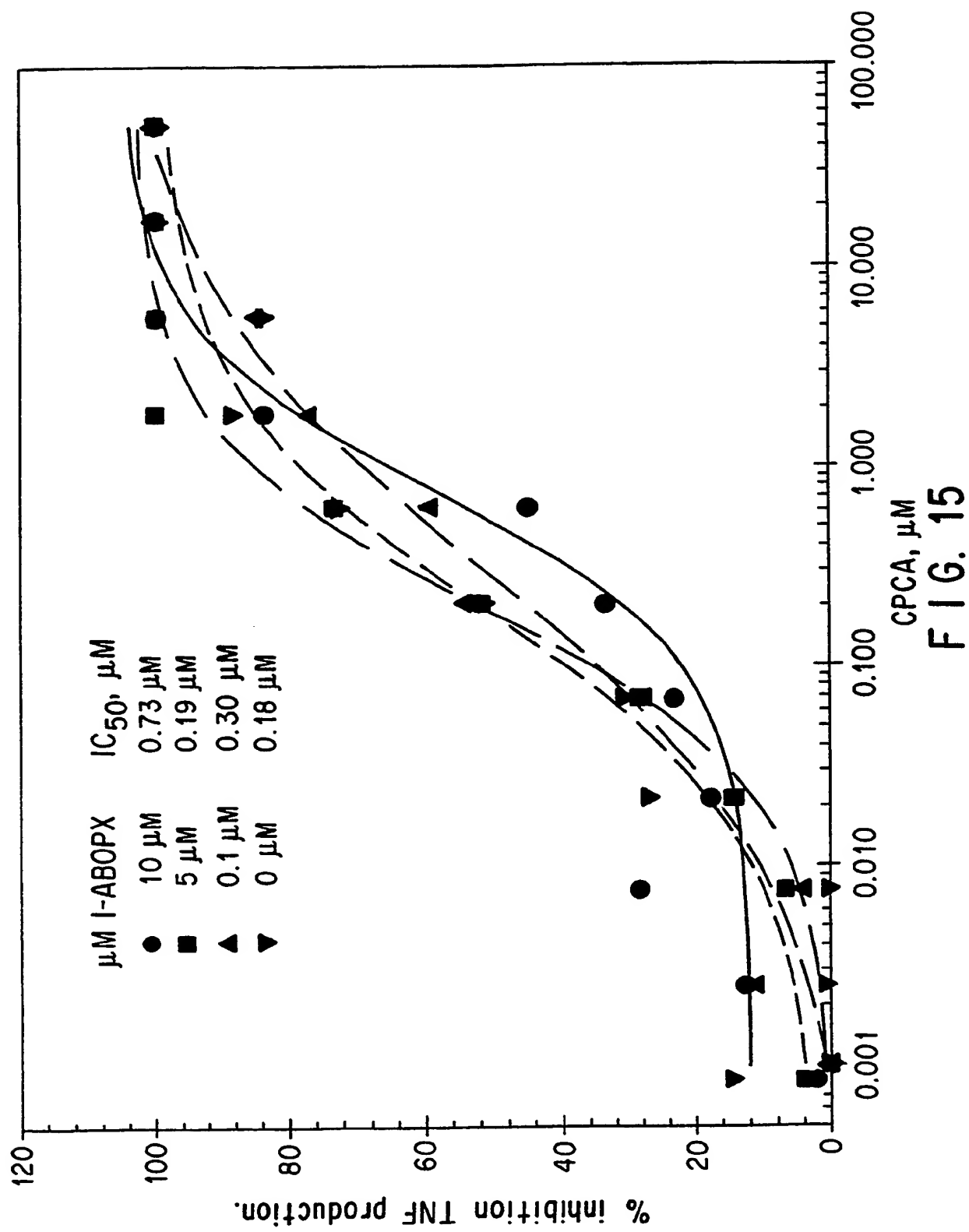


FIG. 13

17/20



18/20



14/22

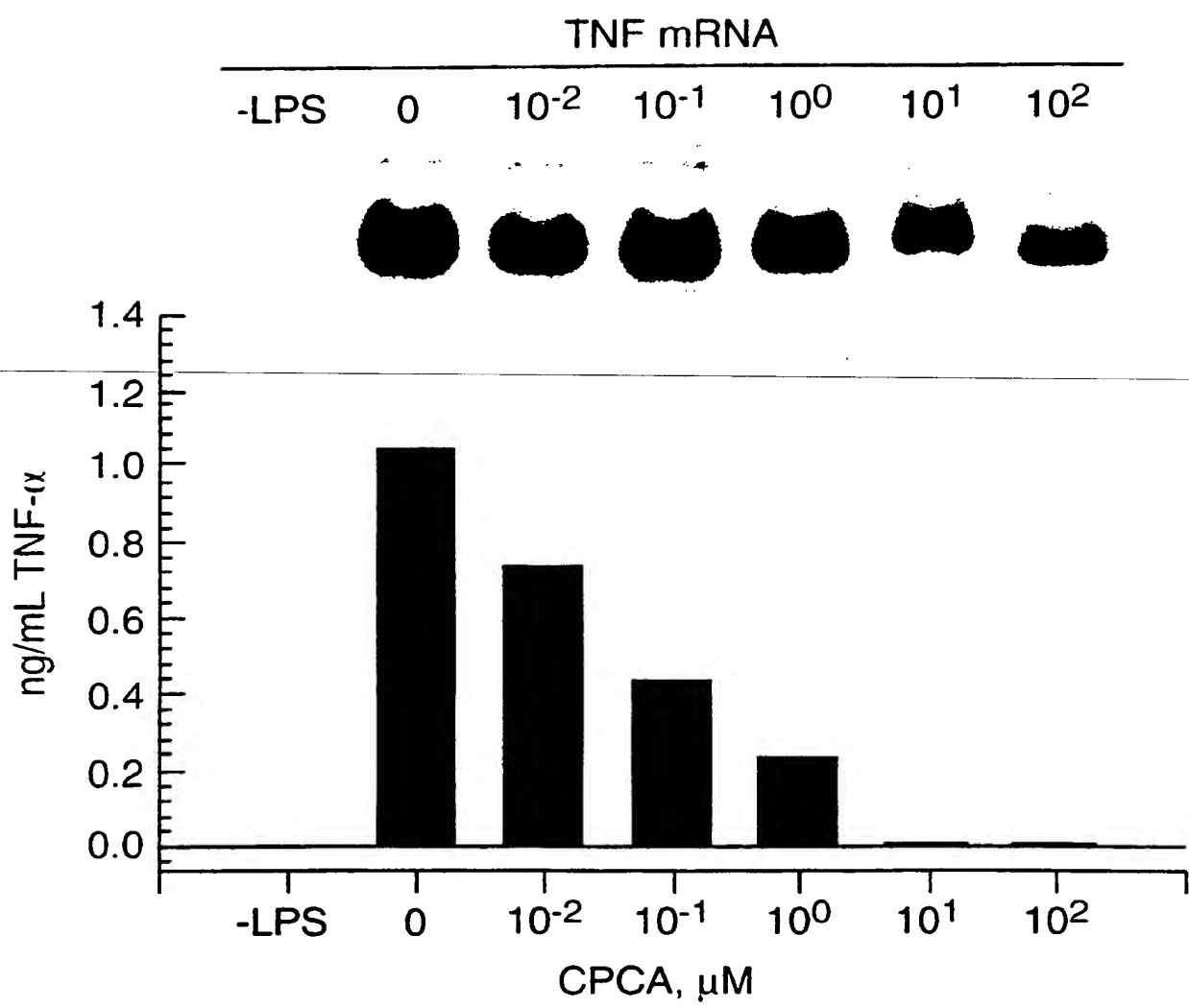


FIG.16

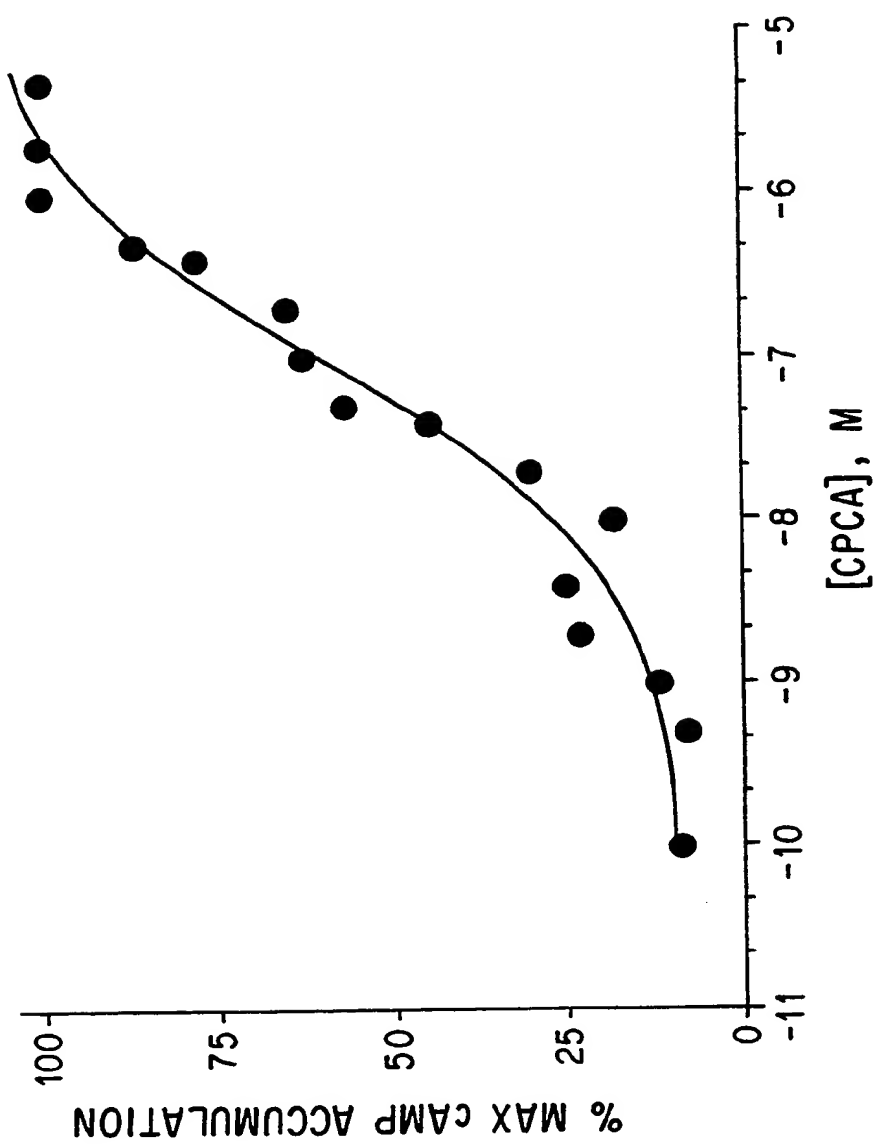


FIG. 17

TITLE OF THE INVENTION

INHIBITION OF TNF $\alpha$  PRODUCTION BY A2b ADENOSINE  
RECEPTOR AGONISTS AND ENHANCERS

5      BACKGROUND OF THE INVENTION

1. FIELD OF THE INVENTION:

The present invention concerns the use of compounds identified as specific modulators of adenosine's physiological actions. The pharmacology of these compounds is characterized through the use of cloned human adenosine A1, A2a, A2b and A3 receptor subtypes. This invention discloses that compounds identified as agonists of the A2b adenosine receptor subtype are useful in inhibiting the production of tumor necrosis factor (TNF $\alpha$ ) by monocytes and/or macrophages. Therefore this invention comprises a method of treatment or prevention of disease states induced by production of TNF $\alpha$ . These conditions include, but are not limited to autoimmune diseases including rheumatoid arthritis, rheumatoid spondylitis, inflammatory bowel disease (ulcerative colitis and Crohn's disease), intestinal pathology associated with graft vs. host disease, organ transplant reactions, septic shock, fever and myalgia due to infection and cachexia associated with chronic infections, malignancy and acquired immune deficiency syndrome, pulmonary diseases such as pulmonary sarcoidosis, silicosis, chronic pulmonary inflammatory disease, adult respiratory distress syndrome.

25     2. BACKGROUND:

Adenosine is a naturally occurring nucleoside which exhibits diverse and potent physiological actions in the cardiovascular, nervous, pulmonary, renal and immune systems. Adenosine has been demonstrated to terminate supraventricular tachycardia through blockage of atrioventricular nodal conduction (J.P. DiMarco, et al., (1985) J. Am. Col. Cardiol. 6:417-425, A. Munoz, et al., (1984) Eur. Heart J. 5:735-738). Adenosine is a potent vasodilator except in the kidney and placenta (R.A. Olsson, (1981) Ann. Rev. Physiol. 43:385-

395). Adenosine produces bronchoconstriction in asthmatics but not in nonasthmatics (Cushly et al., 1984, Am. Rev. Respir. Dis. 129:380-384). Adenosine has been implicated as a preventative agent and in treatment of ventricular dysfunction following episodes of regional or global ischemia (M.B. Forman and C.E. Velasco (1991) Cardiovasc. Drugs and Therapy 5:901-908) and in cerebral ischemia(M.C. Evans, et al., (1987) Neurosci. Lett. 83:287, D.K.J.E.,Von Lubitz, et al., (1988) Stroke 19:1133).

Dog A1 and A2a adenosine receptors were the first adenosine receptors to be cloned. See F. Libert, et al., (1989) Science 244:569-572, C. Maennant, et al., Biochem. Biophys. Res. Comm., (1990) 173:1169-1178, and F. Libert, et al. (1991) EMBO J. 10:1677-1682. The rat A1 adenosine receptor was cloned by L.C. Mahan, et al., (1991) Mol. Pharm. 40:1-7 and S.M. Reppert, et al., (1991) Mol. Endocrin. 5:1037-1048, the rat A2a by Fink et. al., (1992) Mol. Brain Res. 14:186-195, and the rat A2b by Stehle et al. (1992) Mol. Endocrinol. 6:384-393. Cloning of the rat A3 adenosine receptor was reported by Meyerhof et al., (1991) FEBS Lett. 284:155-160 and Zhou et al., (1992) PNAS USA 89:7432-7436. Cloning of the sheep A3 adenosine receptor has been reported by Linden et al., (1993) Mol. Pharm. 44:524-532. Cloning of the human A1, A2a, A2b and A3 receptors were reported in GB 2264948-A (9/15/93). The human A1 adenosine receptor differs by 18 amino acids from the dog A1 sequence and 16 amino acids from the rat A1 sequence. The human A2a adenosine receptor differs by 28 and 71 amino acids, respectively from the dog and rat A2a sequences. The amino acid sequence for the human A3 receptor is 72% identical with the rat A3 receptor and 85% identical with the sheep A3 receptor sequences.

The actions of adenosine are mediated through G-protein coupled receptors, the A1, A2a, A2b and A3 adenosine receptors. The adenosine receptors were initially classified into A1 and A2 subtypes on the basis of pharmacological criteria and coupling to adenylate cyclase (Van Caulker, D., Muller, M. and Hamprecht, B. (1979) J. Neurochem. 33, 999-1003.). Further pharmacological classification of adenosine

receptors prompted subdivision of the A2 class into A2a and A2b subtypes on the basis of high and low affinity, respectively, for adenosine and the agonists NECA and CGS-21680 (Bruns, R.F., Lu, G.H. and Pugsley, T.A. (1986) Mol. Pharmacol. **29**, 331-346; Wan, W., Sutherland, G.R. and Geiger, J.D. (1990) J. Neurochem. **55**, 1763-1771). The existence of A1, A2a and A2b subtypes has been confirmed by cloning and functional characterization of expressed bovine, canine, rat and human receptors. A fourth subtype, A3, had remained pharmacologically undetected until its recent identification by molecular cloning. The rat A3 sequence, tgpcr1, was first cloned from rat testis by Meyerhoff et al. (see above). Subsequently, a cDNA encoding the identical receptor was cloned from striatum and functionally expressed by Zhou et al. (see above). When compared to the other members of the G-protein coupled receptor family, the rat sequence had the highest homology with the adenosine receptors (> 40% overall identity, 58% within the transmembrane regions). When stably expressed in CHO cells, the receptor was found to bind the radioligand  $^{125}\text{I}$ -APNEA ( $\text{N}^6$ -2-(4-amino-3-iodophenyl)ethyladenosine) and when transfected cells were treated with adenosine agonists, cyclic AMP accumulation was inhibited with a potency order of NECA = R-PIA > CGS21680. The rat A3 receptor exhibited a unique pharmacology relative to the A1 and A2 adenosine receptor subtypes and was reported not to bind the xanthine antagonists 1,3-dipropyl-8-phenylxanthine (DPCPX) and xanthine amine congener (XAC). Messenger RNA for the rat A3 adenosine receptor is primarily expressed in the testis.

The sheep homolog of the A3 receptor was cloned from hypophysial pars tuberalis (see Linden et al. above). The sheep receptor is 72% identical to the rat receptor, binds the radioligand  $^{125}\text{I}$ -ABA and is also coupled to inhibition of cyclic AMP. The agonist affinity order of the sheep receptor is I-ABA > APNEA > NECA  $\geq$  R-PIA >> CPA. The pharmacology of xanthine antagonists was extensively studied and the sheep receptor was found to exhibit high affinity for 8-phenylxanthines with para-acidic substitutions. In contrast to the rat transcript, the expression of the sheep A3 adenosine receptor transcript

is widespread throughout the brain and is most abundant in the lung and spleen. Moderate amounts of transcript are also observed in pineal and testis. The cloning and pharmacological profile of the human A3 adenosine receptor was disclosed by Salvatore et al., [P.N.A.S. 90:10365-10369, 1993] and is quite similar to that of the sheep A3 receptor pharmacology.

Based on the use of these cloned receptors, an assay has been described to identify adenosine receptor agonists and antagonists and determine their binding affinity (see GB 2 264 948 A, published 9/15/93; see also R.F. Bruns, et al., (1983) Proc. Natl. Acad. Sci. USA 80:2077-2080; R.F. Bruns, et al.,(1986) Mol. Pharmacol. 29:331-346; M.F. Jarvis, et al. (1989) J. Pharma. Exp. Therap. 251:888-893; K.A. Jacobson et al., (1989) J. Med. Chem. 32:1043-1051).

Adenosine receptor agonists, antagonists and binding enhancers have been identified and implicated for usage in the treatment of physiological complications resulting from cardiovascular, pulmonary, renal and neurological disorders. Adenosine receptor agonists have been identified for use as vasodilators ((1989) FASEB. J. 3(4) Abs 4770 and 4773, (19910 J. Med. Chem. (1988) 34:2570), antihypertensive agents (D.G. Taylor et al., FASEB J. (1988) 2:1799), and anti-psychotic agents (T.G. Heffner et al., (1989) Psychopharmacology 98:31-38). Adenosine receptor agonists have been identified for use in improving renal function (R.D. Murray and P.C. Churchill,(1985) J. Pharmacol. Exp. Therap. 232:189-193). Adenosine receptor allosteric or binding enhancers have shown utility in the treatment of ischemia, seizures or hypoxia of the brain (R.F. Bruns, et al. (1990) Mol. Pharmacol. 38:939-949; C.A. Janusz, et al., (1991) Brain Research 567:181-187). The cardioprotective agent, 5-amino-4-imidazole carboxamide (AICA) ribose has utility in the treatment of ischemic heart conditions, including unstable angina and acute myocardial infarction (H.E. Gruber, et al. (1989) Circulation 80: 1400-1414).

Through the use of homogenous, recombinant adenosine receptors, the identification and evaluation of compounds which have

selectivity for a single receptor subtype is now possible. Because of the variable effects of adenosine documented in other species, the utilization of human adenosine receptor subtypes is advantageous for the development of human therapeutic adenosine receptor agonists, antagonists or enhancers.

5           The anti-inflammatory properties of adenosine have been documented. Adenosine receptor agonists inhibit TNF $\alpha$  production by LPS-stimulated human monocytes (Vraux, et al. 1993 Life Sci. 52:1917-1924) with an affinity profile which does not correspond to A1 or A2a subtype pharmacology. The identification of the specific adenosine receptor subtype mediating the inhibition of TNF $\alpha$  has not been elucidated. With the use of affinity order profiles generated with adenosine receptor agonists, subtype selective adenosine receptor antagonists and information derived from the pharmacological characterization of the human A2b receptor cDNA stably expressed in CHO cells, I have identified the A2b adenosine receptor subtype in mediating the inhibition of TNF $\alpha$  in stimulated human monocytes.

10           The use of an A2b adenosine receptor specific agonist is advantageous over existing therapeutic agents in that a decrease or elimination of side effects experienced when non-selective agonists or the natural agonist, adenosine, are used for therapy. Allosteric effectors or enhancers of the A2b adenosine receptor would eliminate or decrease systemic side effects. Enhancers increase the binding of the native agonists and have been described for A1 adenosine receptors. A2b receptor enhancers remain pharmacologically silent in the absence of adenosine and act locally at sites of inflammation where increases in adenosine concentrations are realized, thereby reducing side effects. The use of such enhancers to inhibit TNF $\alpha$  production naturally forms part of the instant invention.

20           BRIEF DESCRIPTION OF THE DRAWINGS

25           Figure 1   Full length amino acid sequence of human A1 adenosine receptor.

- Figure 2 Full length nucleotide sequence of the cloned human A1 adenosine receptor complementary DNA depicted from the 5' to 3' terminus.
- 5 Figure 3 Full length amino acid sequence of human A2a adenosine receptor.
- 10 Figure 4 Full length nucleotide sequence of cloned human A2a adenosine receptor complementary DNA depicted from the 5' to 3' terminus.
- 15 Figure 5 Full length amino acid sequence of human A2b receptor.
- Figure 6 Full length nucleotide sequence of cloned human A2b adenosine receptor complementary DNA depicted from the 5' to 3' terminus.
- 20 Figure 7 Saturation binding of [<sup>3</sup>H]-cyclohexyladenosine (CHA) to human A1 adenosine receptor in COS7 assay.
- Figure 8 Saturation binding of [<sup>3</sup>H]-CGS21680 to human A2a adenosine receptor in COS7 assay.
- 25 Figure 9 Full length amino acid sequence of human A3 adenosine receptor.
- Figure 10 Full length nucleotide sequence of the cloned human A3 adenosine receptor complementary DNA depicted from the 5' to 3' terminus.
- 30 Figure 11 Adenosine agonists inhibit LPS induced TNF $\alpha$  production in human blood monocytes with a rank order potency of CPC<sub>A</sub>  $\geq$  NECA >> R-PIA > CHA  $\geq$  adenosine > CGS21680. Human peripheral blood mononuclear cells

were cultured on plastic plates coated with fibronectin. The cells were treated with 100 ng/mL of LPS and the indicated concentrations of adenosine agonist. The TNF $\alpha$  levels were measured in cell-culture supernatant by specific ELISA after 18 hours of culture.

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- Figure 12 The adenosine agonist CPCPA inhibits TNF $\alpha$ , but not IL1 $\beta$  or IL-6 release from LPS stimulated human monocytes. Human peripheral blood monocytes were adhered to fibronectin coated plates and stimulated with LPS in the presence of the indicated concentrations of CPCPA. Cell culture supernatant was removed after overnight incubation and tested by specific ELISA for IL-6, IL1 $\beta$ , and TNF $\alpha$ . CPCPA did not inhibit IL-6 or IL1 $\beta$  production.
- Figure 13 The A1 adenosine receptor antagonist DPCPX does not affect the CPCPA induced inhibition of TNF $\alpha$  production in LPS stimulated monocytes. Human peripheral blood monocytes were adhered to fibronectin coated plates. The cells were treated with 100 ng/mL of LPS, and the indicated concentrations of CPCPA and DPCPX. TNF $\alpha$  production levels were measured by specific ELISA in cell-culture supernatant after 18 hours of culture.
- Figure 14 CG21A partially antagonizes CPCPA induced inhibition of TNF $\alpha$  production in LPS stimulated monocytes. Human peripheral blood monocytes were adhered to fibronectin coated plates. The cells were treated with 100 ng/mL of LPS, and the indicated concentrations of CPCPA and CG21A, an adenosine A2a receptor antagonist. TNF $\alpha$  production levels were measured by specific ELISA in cell-culture supernatant after 18 hours of culture. CGS21A inhibited TNF $\alpha$  production in a dose dependent manner in the absence of CPCPA, consistent with the hypothesis that

endogenous adenosine partially represses TNF $\alpha$  production in the assay.

5      Figure 15     The A3 adenosine receptor antagonist I-ABOPX does not affect the CPCA induced inhibition of TNF $\alpha$  production in LPS stimulated monocytes. Human peripheral blood monocytes were adhered to fibronectin coated plates. The cells were treated with 100 ng/mL of LPS, and the indicated concentrations of CPCA and I-ABOPX. TNF $\alpha$  production levels were measured by specific ELISA in cell-culture supernatant after 18 hours of culture.

10     Figure 16    Northern blot analysis of the TNF $\alpha$  mRNA production in LPS stimulated monocytes treated with the adenosine agonist CPCA. Total RNA was extracted from  $1 \times 10^7$  adhered human monocytes one hour following stimulation with LPS in the presence of the indicated concentrations of CPCA. Total RNA (10  $\mu$ g) was blotted using a  $^{32}$ P labeled cDNA probe. No significant reductions in TNF $\alpha$  mRNA production were observed using CPCA at levels sufficient to suppress protein production by greater than ten fold.

15     Figure 17    CPCA dose response of cAMP accumulation in CHO cells stably expressing the human A2b receptor.

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#### SUMMARY OF THE INVENTION

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Adenosine receptor agonists have been shown to inhibit tumor necrosis factor alpha (TNF $\alpha$ ) production in lipopolysaccharide (LPS) stimulated monocytes with an affinity order profile of CPCA  $\geq$  NECA >> R-PIA > CHA  $\geq$  adenosine > CGS21680. This agonist profile does not correlate with either the A1 or A2a adenosine receptor subtype pharmacology. In order to define the receptor subtype mediating the inhibitory effect, adenosine receptor antagonists were

evaluated for their ability to block the inhibition of TNF $\alpha$  production caused by CPCA in LPS-stimulated human monocytes. The involvement of the A1 and A2a adenosine receptor subtypes was ruled out on the basis of the inability of DPCPX and 3-succinylaminostrylcaffine, CG21A, respectively, to appreciably antagonize the inhibition produced by CPCA. The A3 adenosine receptor specific antagonist IABOPX was also ineffective in blocking agonist induced inhibition of TNF $\alpha$  production. The agonist affinity order profile established for the monocyte adenosine receptor was similar to the A2b receptor in VA13 human fibroblasts and human erythroleukemic cells (HEL) defined by EC50 values for intracellular cyclic adenosine monophosphate (cAMP) accumulation. However, the potency of the agonists to inhibit TNF $\alpha$  production in monocytes was greater than values determined by increases in cAMP accumulation in fibroblasts or HEL cells. I have found that in stable CHO cells expressing the cloned human A2b cDNA, the potency (EC50) of CPCA to induce cAMP accumulation was similar to the value obtained for inhibition of TNF $\alpha$  production in LPS-stimulated human monocytes. To define which adenosine receptor subtypes are present on monocytes, A1, A2a, A2b, and A3 adenosine receptor transcripts were detected by reverse transcriptase PCR (RT-PCR) of mRNA prepared from both LPS-stimulated and non-stimulated monocytes. The regulation of TNF $\alpha$  expression resulting from mediation at the A2b receptors is demonstrated to be consistent with a mechanism involving increased intracellular cAMP levels.

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ABBREVIATIONS

[<sup>3</sup>H]-CHA, [<sup>3</sup>H]-cyclohexyladenosine; [<sup>3</sup>H]-NECA, [<sup>3</sup>H]-5'-N-ethyl-carboxamido-adenosine; <sup>125</sup>I-ABA, N<sup>6</sup>-(4-amino-3-  
125iodobenzyl)adenosine; <sup>125</sup>I-APNEA, N<sup>6</sup>-2-(4-amino-3-  
125iodophenyl)ethyladenosine; NECA, 5'-N-  
5 ethylcarboxamidoadenosine; CGS21680, 2-[4-(2-  
carboxyethyl)phenyl]ethylamino-5'-N-ethylcarboxamidoadenosine;  
(R,S)-PIA, (R,S)-N<sup>6</sup>-phenyl-2-propyladenosine; CPA, N<sup>6</sup>-  
10 cyclopentyladenosine; CPCPA, 5'-(N-cyclopropyl)-  
carboxamidoadenosine; CG21A, 3-succinylaminostrylcaffine; I-  
ABOPX, (3-(3-iodo-4-aminobenzyl)-8-(4-oxyacetate)phenyl-1-  
15 propylxanthine; BW-A1433, 1,3-dipropyl-8-(4-  
acrylate)phenylxanthine; XAC, xanthine amine cogener; DPCPX, 1,3-  
dipropyl-8-cyclopentylxanthine; GTP $\gamma$ S, guanosine 5'-O-3-  
thiotriphosphate; Gpp(NH)p, 5'-guanylimidodiphosphate; G protein,  
15 guanine nucleotide-binding proteins.

DETAILED DESCRIPTION OF THE INVENTION

This invention provides a method for achieving specific inhibition of TNF $\alpha$  production through agonist stimulation of the A2b receptor. TNF $\alpha$  is a pro-inflammatory cytokine which, among other effects, induces fever and stimulates phospholipase A2 production. Lipopolysaccharide (LPS) is a biological mediator which gives rise to a number of adverse responses. A principal mediator to these effects is TNF $\alpha$ . A variety of adenosine receptor agonists have been tested for their ability to block LPS-mediated TNF $\alpha$  production in human monocytes [Le Vraux et al., Life Sciences 52:1917-1924, 1993]. Figure 11 summarizes the pharmacological profile of this effect [CPCA  $\geq$  NECA >> R-PIA > CHA  $\geq$  adenosine > CGS21680]. The conclusion reported in Le Vraux et al., based on this pharmacology, was that the inhibition of TNF $\alpha$  production was probably mediated through the A3 adenosine receptor, or through an uncharacterized receptor, but not through the A1 or A2 adenosine receptors. As can be seen from this data, CPCPA and NECA are the most potent inhibitors of TNF $\alpha$ .

production. Both compounds have been characterized as binding both the A1 and the A2 adenosine receptor subtypes with high affinity, see the table below:

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AFFINITY OF ADENOSINE ANALOGS FOR HUMAN ADENOSINE  
RECEPTOR SUBTYPES,  $K_i$  or  $K_d$ ,  $\mu M$  \*

| Agonists | A1          | A2a         | A2b          | A3        |       |
|----------|-------------|-------------|--------------|-----------|-------|
| 5        | NECA        | 0.025       | 0.029        | 0.9 (a)   | 0.026 |
|          | CPCA        | 0.006 (rat) | 0.0134 (rat) | 0.050 (a) | 1.0   |
|          | CGS21680    | 56          | 0.017        | 1600 (b)  | 5.6   |
|          | R-PIA       | 0.003       | 0.127        | 160 (b)   | 0.034 |
|          | CHA         | 0.002       | 0.6          | 280 (b)   | n.d.  |
| 10       | Antagonists |             |              |           |       |
|          | DPCPX       | 0.0007      | 0.10         | 0.55 (b)  | 0.75  |
|          | CGS21A      | 35 (rat)    | 0.143 (rat)  | n.d.      | >50   |

15 \*Values determined in rat are indicated, otherwise all other data is from human, (a) EC<sub>50</sub> values for cAMP accumulation in stable CHO cells expressing the human A2b cDNA; (b) EC<sub>50</sub> values for cAMP accumulation in human erythroleukemic cells, HEL cells.

20 The A1 adenosine receptor selective agonists R-PIA and CHA are significantly less potent than CPCA or NECA. The A2a specific agonist CGS21680 was found to be the least potent of all. The rank order of potency of the compounds to inhibit TNF $\alpha$  production is not like that of either A1 or A2 [Le Vraux et al., Life Sciences 52:1917-1924, 1993]. The affinity order profile reported by Le Vraux et al. is similar to the agonist profile reported by Castanon and Spevak [BBRC 198:626-631, 1994] for the induction of cyclic adenosine monophosphate (cAMP) accumulation in stable CHO cell lines expressing the cloned A2b adenosine receptor. However, Castanon and Spevak did not study the role of the A2b receptor in inhibition of TNF $\alpha$  production. In addition, the agonist affinity order profile data reported by Le Vraux et al. for TNF $\alpha$  inhibition is not dissimilar from the agonist order profile reported by Salvatore et al., [P.N.A.S. 90:10365-10369, 1993] for the cloned A3 adenosine receptor and suggested that

the A3 receptor may be responsible for TNF $\alpha$  inhibition in LPS-stimulated monocytes. However, the potency of CPCPA for the A3 receptor was not reported by Salvatore et al. and therefore, prior to this invention, the role of A3 adenosine receptor in the inhibition of TNF $\alpha$  production could not be ruled out and the specific adenosine receptor subtype which is responsible for inhibition of TNF $\alpha$  production could not be positively identified. This patent disclosure demonstrates that CPCPA has a much lower affinity for the A3 receptor than it does for the A2b receptor and by using A3 adenosine receptor specific antagonists, the involvement of A3 receptor activation in the inhibition of TNF $\alpha$  production is definitively ruled out. This patent disclosure demonstrates that A1 and A2a adenosine receptors are not involved in the inhibition of TNF $\alpha$  production. This invention reveals that activation only at the A2b adenosine receptor is responsible for the inhibition of TNF $\alpha$  production.

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The role of cAMP elevations has been correlated with the inhibition of LPS induced TNF $\alpha$  production defined through the use of the phosphodiesterase inhibitor pentoxifyllin [Strieter, et al., (1988) Biochem. Biophys. Res. Commun. 155: 1230-1236]. The inhibition of TNF $\alpha$  production through activation at A2b adenosine receptors on stimulated monocytes is therefore consistent with a mechanism resulting from increases in intracellular cAMP. Therefore, this invention comprises a method for inhibiting TNF $\alpha$  production specifically through A2b receptor activation.

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Since Le Vraux et al., suggested that the receptor responsible for inhibition of TNF $\alpha$  production was possibly the A3 adenosine receptor and not the A1 or A2 receptors, I initiated the following studies in order to elucidate which receptor is, in fact, responsible for inhibition of TNF $\alpha$  production.

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I confirmed that the A1 and A2a receptor subtypes are not responsible for the inhibition of TNF $\alpha$  production by using the A1 and

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A<sub>2a</sub> adenosine receptor selective antagonists DPCPX and CG21A respectively. These compounds do not appreciably alter the IC<sub>50</sub> of CPC<sub>A</sub> in antagonist competition experiments except at very high concentrations (see Figures 13 and 14). This data confirms that the A<sub>1</sub> and A<sub>2a</sub> adenosine receptor subtypes are not involved in the inhibition of TNF $\alpha$  production. I confirmed that the A<sub>3</sub> receptor subtype was not responsible for the inhibition of TNF $\alpha$  production by using the A<sub>3</sub> specific antagonist, I-ABOPX (Figure 15). I-ABOPX did not alter the IC<sub>50</sub> of CPC<sub>A</sub> inhibition of TNF $\alpha$  production.

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I further determined that the affinity of CPC<sub>A</sub> for the A<sub>3</sub> adenosine receptor subtype is 1  $\mu$ M and therefore, the A<sub>3</sub> receptor cannot be responsible for the inhibition of TNF $\alpha$  production induced by CPC<sub>A</sub> which exhibits a much higher (20,000-fold) affinity for the A<sub>2b</sub> than the A<sub>3</sub> adenosine receptor. I obtained the EC<sub>50</sub> value for CPC<sub>A</sub> induced cAMP accumulation in stable CHO cell lines expressing the human A<sub>2b</sub> receptor and found that the EC<sub>50</sub> value is the same as that obtained from the stimulated monocytes (Figure 17). I further confirmed that the effect is specific for TNF $\alpha$  because IL1 $\beta$  and IL-6 production are unaffected by treatment with CPC<sub>A</sub>, (Figure 12).

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Northern blot data of total RNA from LPS stimulated monocytes indicates that titration of CPC<sub>A</sub> reduces the levels of secreted TNF $\alpha$  protein in a dose dependent manner, Figure 16. This data indicates that adenosine agonists inhibit TNF $\alpha$  production primarily through post-transcriptional mechanisms. This observation is consistent with reports that TNF $\alpha$  mRNA contains 3'-untranslated sequences that mediate translational activation in response to specific inducing signals (e.g. LPS). Removal of these sequences has been shown to result in the inability of the mRNA to be translated. Therefore, it appears that adenosine blocks components of the LPS signal transduction pathway that are related to these 3'-untranslated elements of the TNF $\alpha$  gene.

To define which adenosine receptor subtypes are present on monocytes, A<sub>1</sub>, A<sub>2a</sub>, A<sub>2b</sub>, and A<sub>3</sub> adenosine receptor transcripts were detected by reverse transcriptase PCR (RT-PCR) of mRNA prepared from both LPS-stimulated and non-stimulated monocytes. All four

adenosine receptor subtypes were detected in mRNA prepared from both normal and LPS-stimulated monocytes. Even though all of the identified adenosine receptor subtypes are present on monocytes, this invention reveals that only the A2b receptor affects TNF $\alpha$  production.

5 Therefore, one embodiment of this invention is a method for identifying A2b adenosine receptor selective compounds which comprises the steps of:

- (a) contacting monocytes with a test compound and measuring the effect of the test compound on TNF $\alpha$  production;
- 10 (b) contacting a test compound, identified according to step (a) as inhibiting TNF $\alpha$  production by the monocytes, with membranes derived from a stable cell line individually expressing each of the A1, A2a, A2b, or A3 adenosine receptor or with the whole cell individually expressing each of the A1, A2a, A2b, or A3 adenosine receptor and measuring the binding affinity of the test compound for the receptor or the effect of the test compound on cAMP production in the stable cell line;
- 15 (c) selecting compounds which bind to the A2b adenosine receptor or which induces elevation in cAMP in the cell line expressing the A2b adenosine receptor and which do not bind to membranes or affect the cAMP level in the stable cell lines expressing the A1, A2a, or A3 adenosine receptor subtypes.

20 This invention likewise comprises the use of compounds identified according to this method which have A2b adenosine receptor enhancer or agonist activities for the inhibition of TNF $\alpha$  production. This invention further comprises a method for inhibiting TNF $\alpha$  production by contacting monocytes with inhibitorily effective amounts of compounds that act as A2b adenosine receptor agonists. An inhibitorily effective amount of an A2b adenosine receptor agonist is, for example, 0.1 ng to 10 mg/kg per day of CPCPA, NECA or a compound exhibiting similarly potent or more potent A2b adenosine receptor agonist properties.

The following examples are provided to further define but not to limit the invention defined by the foregoing description and the claims which follow:

EXAMPLE 1

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STEP A:

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In the first step of obtaining the partial cDNAs encoding the human A1 and A2a adenosine receptors, total RNA was extracted by homogenizing 2.3g human ventricle in 20 ml 5M guanidine isothiocyanate, 0.1M sodium citrate, pH 6.3, 1mM EDTA, pH 7.0, 5% beta-mercaptoethanol, and 0.5% sodium lauryl sarcosinate. The homogenate was centrifuged for 10 min. at 10,000 rpm and the resulting supernatant was layered onto a cushion of 5.7M CsCl/0.1M EDTA, pH 7.0. After 20 hrs. of centrifugation at 24,000 rpm, the resulting pellet was precipitated one time and then passed over an oligo(dT)-cellulose (PHARMACIA, Piscataway, NJ) column to isolate poly(A)+ RNA.

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An oligo(dT) primed library was synthesized from 5 µg of the poly(A)<sup>+</sup> human ventricle RNA using the YOU-PRIME cDNA SYNTHESIS KIT (PHARMACIA, Piscataway, NJ). See Gubler and Hoffman Gene 25:263 (1983). The resulting double-stranded cDNA was ligated into λgt10 EcoRI arms (PROMEGA, Madison, WI) and packaged according to the GIGAPACK II GOLD PACKAGING EXTRACT protocol (STRATAGENE, La Jolla, CA). See Huynh et al. (1985) DNA Cloning Techniques: A Practical Approach, IRL Press, Oxford, p.49 and Kretz et al. Res. 17:5409.

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The E. coli strain C600Hfl (PROMEGA, Madison, WI) was infected with library phage, plated on agar plates, and incubated at 37°C. The phage DNA was transferred to HYBOND-N nylon membranes (AMERSHAM, Arlington Heights, IL) according to the manufacturer's specifications.

Synthetic probes were constructed from overlapping oligonucleotides (A1 probe: 62+63, A2 probe: 52+53; see Table I for their sequences) based on the published dog A1 (RDC7 ) and

A2a(RDC8) sequences (F Libert, et al,(1989) Science 244:569-572).  
The oligonucleotides were annealed and filled-in with a<sup>32</sup>P-dCTP  
(NEN, Wilmington, DE) and Klenow enzyme. The filters were  
hybridized with the appropriate probe in 5XSSC, 30% formamide,  
5XDenhardt's solution, 0.1% SDS, and 0.1mg/ml sonicated salmon  
5 sperm DNA at 42°C, overnight. Following hybridization the filters  
were washed to a final stringency of 6XSSC at 50°C and exposed to X-  
OMAT AR film (KODAK, Rochester, NY) at -70°C. The resulting  
positives were plaque purified by two additional rounds of plating and  
hybridization. Insert DNA was excised with NotI and ligated into NotI  
10 digested pBLUESCRIPT II KS+ (STRATAGENE, La Jolla, CA).  
(Genebank # 52327) DNA sequences were determined by the  
SEQUENASE protocol (USBC, Cleveland, OH). See Tabor and  
Richardsaon, J. Biol. Chem. 264 pp 6447-6458. Two clones were  
isolated in these screens. The human ventricle A1 cDNA (hva1-3a) and  
15 human ventricle A2a cDNA (hva2-13) contain portions of coding  
sequences for proteins homologous to the reported dog A1 and A2a  
cDNAs, respectively. The coding region of the human A1 clone  
corresponds to nucleotides 482 through 981 (Figure 2) and is 92%  
identical to the dog A1 sequence at the nucleotide level. The coding  
region of the human A2a clone corresponds to nucleotides 497 through  
20 1239 (Figure 4), and is 90% identical to the dog A2a sequence at the  
nucleotide level.

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STEP B:

The human ventricle A1 adenosine receptor partial cDNA (hvA1-3a) is a 543 bp NotI fragment containing 23 bp 3' untranslated sequence and is 460 bp short of the initiation methionine based on sequence homology to the dog A1 cDNA. A modification of the 5' RACE (rapid amplification of cDNA ends) method (MA Frohman et al,(1988), Proc. Natl. Acad. Sci. USA, 85:8998-9002) was used to generate the 5' coding region of the cDNA. First strand cDNA was synthesized from 1 $\mu$ g of the human ventricle poly(A)<sup>+</sup> RNA in a total volume of 40ml containing 50mM Tris, pH 8.0, 140mM KCl, 10mM MgCl<sub>2</sub>, 10mM DTT, 15mM each dNTP, 20 units RNasin (PROMEGA, Madison, WI), 20pmol human primer 79 (see Table I), and 9.2 units AMV reverse transcriptase at 37°C for 2 hrs. The reaction was then diluted to 120  $\mu$ l with 0.5 mM Tris, pH 7.6/0.05 mM EDTA and passed through a SEPHACRYL S-300 SPUN COLUMN (PHARMACIA, Piscataway, NJ). The product in the column effluent was polyadenylated in 100mM potassium cacodylate, pH 7.2, 2mM CoCl<sub>2</sub>, 0.2mM DTT, 0.15mM dATP, and 14 units terminal deoxynucleotidyl transferase in a total volume of 31 $\mu$ l for 10 min. at 37°C. The reaction was terminated by heating at 65°C for 15 min. and then diluted to 500 ml with 10 mM Tris, pH 8.0/1 mM EDTA (TE).

Ten  $\mu$ l of the poly(A)-tailed first strand cDNA was used as template in a primary PCR amplification reaction according to the GENEAMP protocol (PERKIN ELMER CETUS, Norwalk, CT; see Saiki et al. (1988) Science 239:487-491) containing 10pmol primer 70, 25pmol primer 71, and 25pmol human primer 80 (see table I) in a total volume of 50 ml. Primer 70 is 5'-gactcgagtgcacatcga(t)<sub>17</sub>, primer 71 is 5'-gactcgagtgcacatcga, and both are based on MA Frohman, et al (1988), Proc. Natl. Acad. Sci. USA, 85:8998-9002. One cycle of PCR was performed of 1 min at 95°C, 2 min at 50°C, 40 min at 72°C, followed by 40 cycles of 40 sec at 94°C, 2 min at 56°C, 3 min at 72°C. The primary PCR amplification reaction product was electrophoresed through a 1.4% agarose gel and an area corresponding to approximately 600 bp was excised. The gel slice was melted and 1  $\mu$ l was used as

template in a secondary PCR amplification reaction containing 100pmol primer 71 and human primer 81 (see Table I) for 30 cycles of 1 min at 94°C, 2 min at 56°C, 3 min at 72°C. The secondary PCR amplification product was digested with EcoRI and SalI and electrophoresed on a 1.4% agarose gel. An area corresponding to 500-600bp was excised and ligated into EcoRI/SalI digested pBLUESCRIPT II KS+ (STRATAGENE, La Jolla, CA). The sequence of the 515 bp PCR product (5'HVA1-9) was determined by the SEQUENASE protocol (USBC, Cleveland, OH). The partial human ventricle A1 cDNA and the PCR product contain overlapping sequence and represent the complete coding region for the human A1 receptor, including 14 and 23 bp of 5' and 3' untranslated sequences, respectively. The sequence of the human A1 adenosine receptor cDNA so identified, is shown in Figure 2.

15 STEP C:

A probe was generated by Klenow enzyme extension, including a<sup>32</sup>P-dCTP, of annealed oligonucleotides 62 and 63, and used to screen a human kidney cDNA library (CLONTECH, Palo Alto, CA). E. coli strain C600hfl (PROMEGA, Madison, WI) was infected with library phage and grown overnight on agar plates at 37°C. Phage DNA was transferred to HYBOND-N nylon filters according to the manufacturer's protocol (AMERSHAM, Arlington Heights, IL). The probe was incubated with the filters in 750mM NaCl, 75mM sodium citrate, 30% formamide, 0.1% sodium dodecyl sulfate, 0.5mg/mL polyvinylpyrrolidone, 0.5mg/mL bovine serum albumin, 0.5mg/mL Ficoll 400, and 0.1mg/mL salmon sperm DNA, at 42°C overnight. The filters were washed in 0.9M NaCl and 90mM sodium citrate at 50°C. A positively hybridizing phage (hkA1-14), was identified and purified by replating and screening with the probe twice more. The final phage plaque was transferred to 0.5 mL 50mM Tris, pH 7.5, 8mM MgSO<sub>4</sub>, 85 mM NaCl, 1mg/mL gelatin, and 1 μL of a 1:50 dilution in water of the phage stock was used as template for PCR amplification. 50 pmol each of 1amL and 1amR (Table I) oligonucleotide primers were included, and subjected to 30 cycles of 40 sec at 94°C, 1 min at 55°, 3 min at 72°,

then a final 15 min at 72°, according to the GENEAMP protocol (PERKIN ELMER CETUS, Norwalk, CT). A 2.0 kb product was identified by agarose gel electrophoresis, and this was subcloned into the EcoRI site of pBLUESCRIPT II KS+ (STRATAGENE, La Jolla, CA). Sequence analysis by the SEQUENASE protocol (USBC, Cleveland, OH) demonstrated that this cDNA was homologous to the reported dog A1 clone. SmaI and EcoRI digestion released a DNA fragment containing coding sequence from base pair 76 through the translation STOP codon (Figure 2) that is identical to the human ventricle A1 cDNA sequence (clones hva1-3a and 5'hva1-9). This fragment was used in construction of the full length coding sequence (see below). The human kidney cDNA also includes about 900 bp of 3' untranslated sequence.

15 STEP D:

The human ventricle A2a adenosine receptor partial cDNA (hvA2-13) is a 1.6 kb NotI fragment containing approximately 900 bp 3' untranslated sequence and is 496 bp short of the initiation methionine based on sequence homology to the dog A2a cDNA clone. Two consecutive rounds of 5' RACE were utilized to generate the 5' coding region of the cDNA. First strand cDNA was synthesized from 1 µg of the human ventricle poly(A)<sup>+</sup> RNA in a total volume of 40 ml containing 50mM Tris, pH 8.0, 140mM KCl, 10mM MgCl<sub>2</sub>, 10mM DTT, 15mM each dNTP, 20 units RNasin, 20pmol human primer 68 or 74 (for 1st or 2nd round RACE respectively), and 9.2 units AMV reverse transcriptase at 37°C for 2 hrs. The reaction was then diluted to 120ml with 0.5 mM Tris, pH 7.6/0.05 mM EDTA and passed through a SEPHACRYL S-300 SPUN COLUMN. The products in the column effluents were polyadenylated in 100mM potassium cacodylate, pH 7.2, 2 mM CoCl<sub>2</sub>, 0.2 mM DTT, 0.15 mM dATP, and 14 units terminal deoxynucleotidyl transferase in a total volume of 31 µl for 10 min. at 37°C. The poly(A) tailing reaction was terminated by heating at 65°C for 15 min. and then diluted to 500 µl with TE.

Five or 10  $\mu$ l (for 1st or 2nd round RACE respectively) of the poly(A) tailed first strand cDNA was used as template in the PCR amplification reaction according to the GENEAMP protocol containing 10pmol primer 70, 25 pmol primer 71 (primer 70 and 71 sequences are given above), and 25 pmol human primer 69 or 75 (1st or 2nd round RACE respectively; see Table I) in a total volume of 50  $\mu$ l. One cycle of PCR was performed of 1 min at 95°C, 2 min at 50°C, 40 min at 72°C, followed by 40 cycles of 40 sec at 94°C, 2 min at 56°C, 3 min at 72°C. The PCR amplification products were digested with EcoRI and Sall and electrophoresed on a 1.4% agarose gel. Areas corresponding to 200-400 bp were excised and ligated into EcoRI/Sall digested pBLUESCRIPT II KS+ (STRATAGENE, La Jolla, CA). The sequences of the two A2a PCR products, the 332 bp 1st round RACE product (5'hvA2-14) and the 275 bp 2nd round RACE product (5'hva2-29) were determined by the SEQUENASE (USBC, Cleveland, OH) protocol. By sequence homology comparisons with the dog A2a adenosine receptor cDNA sequence, the 1st round RACE product (5'hvA2-14) was 258 bp short of the initiation methionine and the second round RACE product (5'HVA2-29) was determined to extend 1bp upstream of the initiation methionine. The human ventricle A2a partial cDNA clone (hvA2-13) and the human A2a PCR products (5'hvA2-14 and 5'hva2-29) contain overlapping sequence and together represent the complete coding sequence for the human adenosine A2a receptor, and include 1 bp and 0.8 kb of 5' and 3' untranslated sequence, respectively. The sequence of the human A2a adenosine receptor is shown in Figure 4.

STEP E:

A double-stranded DNA probe was generated by Klenow enzyme extension, including a<sup>32</sup>P-dCTP, of annealed oligonucleotides 66 and 67, and used to screen a human striata cDNA library (STRATAGENE, La Jolla, CA). The oligonucleotide sequence was based on a region of the human ventricle A2a cDNA sequence. E. coli strain XL1-blue (STRATAGENE, La Jolla, CA) cells were infected with library phage and grown overnight on agar plates at 37°C. Phage DNA was transferred to HYBOND-N nylon filters according to the manufacturer's protocol (AMERSHAM, Arlington Heights, IL). The probe was incubated with the filters in 750 mM NaCl, 75 mM sodium citrate, 10% formamide, 0.5% sodium dodecyl sulfate, 0.5 mg/mL polyvinylpyrrolidone, 0.5 mg/mL bovine serum albumin, 0.5 mg/mL Ficoll 400, and 0.02 mg/mL salmon sperm DNA, at 42°C overnight. The filters were washed in 0.9 M NaCl and 90 mM sodium citrate at 50°C. A positively hybridizing phage (hbA2-22A) was identified and purified by replating and screening with the probe twice more, and subcloned into the plasmid pBLUESCRIPT SK- by the manufacturer's protocol (STRATAGENE, La Jolla, CA). See Short et al. (1988) Nucl. Acids Res. 16:7583-7600; Sorge (1988) Stratagies 1:3-7. The human brain A2a adenosine receptor cDNA (hbA2-22A) spans bp 43 of the A2 coding sequence (Figure 4) through the translation STOP codon, and includes about 900 bp of 3' untranslated sequence. The sequence of this human brain A2a cDNA is identical to the human ventricle A2a adenosine receptor cDNA (hvA2-13, 5'hvA2-14 and 5'hvA2-29).

STEP F:

A double-stranded DNA probe was generated by Klenow enzyme extension of annealed oligonucleotides 129 and 130, including a<sup>32</sup>P-dCTP, and used to screen a human frontal cortex cDNA library (STRATAGENE, La Jolla, CA). The oligonucleotide sequence was based on a region of the human A2a and A1 cDNA sequence. E. coli strain XL-1 blue (STRATAGENE, La Jolla, CA) cells were infected with library phage and grown overnight at 37°C. Phage DNA was

transferred to HYBOND-N nylon filters according to the manufacturer's protocol (AMERSHAM, Arlington Heights, IL). The probe was incubated with the filters in 750 mM NaCl, 75 mM sodium citrate, 10% formamide, 0.5% sodium dodecyl sulfate, 0.5 mg/mL polyvinyl-pyrrolidone, 0.5 mg/mL bovine serum albumin, 0.5 mg/mL Ficoll 400, and 0.02 mg/mL salmon sperm DNA, at 42°C overnight.  
5 The filters were washed in 0.9 M NaCl and 90 mM sodium citrate at 50°C. A positively hybridizing phage (hb-32c), was identified and purified by replating and screening with the probe twice more. The insert was subcloned to the plasmid pBLUESCRIPT SK- according to the manufacturer's protocol (STRATAGENE, La Jolla, CA). Sequence analysis by the SEQUENASE protocol (USBC, Cleveland, OH) demonstrated a complete open reading frame coding for amino acid sequence homologous to both of the previously isolated human A1 and A2a clones. This homologous adenosine receptor subtype-cDNA is the  
10 A2b subtype having the sequences in Figures 5 and 6. A 1.3 kb SmaI-XmnI fragment was ligated into the SmaI site of pSVL (PHARMACIA, Piscataway, NJ), giving the full length coding sequence of the A2b adenosine receptor in a plasmid suitable for its expression in COS and CHO cells. See Sprague et al. (1983) *J. Virology* 45:773; Templeton and Eckhart (1984) *Mol. Cell Biol.* 4:817.  
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Table I:

Sequences and directions of the primers used in the isolation of cDNA's and construction of expression plasmids, along with the positions in the clones upon which the sequences are based. Dog A1 and A2a cDNA clones are from F. Libert, et al, (1989) *Science* 244:569-572. Primers LamL and LamR are based on the sequence of λgt10 (T.V. Hyunh, et al. (1985) *DNA Cloning: A Practical Approach*, Vol 1, D. Glover, ed, IRL Press, Oxford). The A2b adenosine receptor subtype encoded by the clone hb32C was determined to be the A2b adenosine receptor subtype on the basis of the binding profile of the adenosine receptor agonist NECA and affinities for adenosine receptor  
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antagonists measured on membranes prepared from pSVLhb32C  
transfected COS7, CHO or HEK 293 cells.

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|    |    | name                    | sequence  | position  | clone     | direction |
|----|----|-------------------------|-----------|-----------|-----------|-----------|
| 5  | 52 | ATTCGCAGCCACGTCTGA-     | 1201-1260 | dog A2a   | sense     |           |
|    |    | GGCGGCGGGAGCCCTCAA-     |           |           |           |           |
|    |    | AGCAGGTGGCACCAAGTGCC-   |           |           |           |           |
|    |    | CGC (SEQ ID NO. 1)      |           |           |           |           |
| 10 | 53 | GCGGAGGGCTGATCTGCT-     | 1305-1246 | dog A2a   | antisense |           |
|    |    | CTCCCATCACTGCCATGAG-    |           |           |           |           |
|    |    | CTGCCAAGGCGCGGGCAC-     |           |           |           |           |
|    |    | TGGTGCC (SEQ. ID NO. 2) |           |           |           |           |
| 15 | 62 | TCCAGAACAGTTCCGGGTCA-   | 958-1017  | dog A1    | sense     |           |
|    |    | CCTTCCTTAAGATCTGGAA-    |           |           |           |           |
|    |    | TGACCAACTCCGCTGCCAGC-   |           |           |           |           |
|    |    | CCA (SEQ. ID NO. 3)     |           |           |           |           |
| 20 | 63 | AGTCGTGGGGCGCCTCCT-     | 1062-1003 | dog A1    | antisense |           |
|    |    | CTGGGGGGTCCTCGTCGAC-    |           |           |           |           |
|    |    | GGGGGGCGTGGGCTGGCAG-    |           |           |           |           |
|    |    | CGGA (SEQ ID NO. 4)     |           |           |           |           |
| 25 | 66 | GCCTCTTGAGGATGTGG-      | 500-542   | 5'hvA2-14 | sense     |           |
|    |    | TCCCCATGAACATACATGGT-   |           |           |           |           |
|    |    | GTACTTCA (SEQ ID NO. 5) |           |           |           |           |
|    | 67 | GCAGGGGCACCAGCACACA-    | 572-528   | 5'hva2-14 | antisense |           |
| 30 |    | GGCAAAGAAGTTGAAGTAC-    |           |           |           |           |
|    |    | ACCATGT (SEQ ID NO. 6)  |           |           |           |           |

|    | name | sequence  | position | clone     | direction |
|----|------|---|----------|-----------|-----------|
| 5  | 68   | TCGCGCCGCCAGGAAGAT<br>(SEQ ID NO 7)   | 616-599  | hva2-13   | antisense |
|    | 69   | TATATTGAATTCTAGACAC-<br>CCAGCATGAGC (SEQ ID NO.8)                             | 591-574  | hva2-13   | antisense |
| 10 | 74   | TCAATGGCGATGGCCAGG<br>(SEQ ID NO. 9)  | 303-286  | 5'hva2-14 | antisense |
| 15 | 75   | TATATTGAATTCATGGA-<br>GCTCTGCGTGAGG-<br>(SEQ ID NO. 10)                       | 276-259  | 5'hva2-14 | antisense |
|    | 79   | GTAGACCATGTACTCCAT<br>(SEQ ID NO. 11)   | 560-543  | hva1-3a   | antisense |
| 20 | 80   | TATATTGAATTCTGACCT-<br>TCTCGAACTCGC-<br>(SEQ ID NO. 12)                       | 537-521  | hva1-3a   | antisense |
| 25 | 81   | ATTGAATTGATCACGGG-<br>CTCCCCCATGC-<br>(SEQ ID NO. 13)                         | 515-496  | hva1-3a   | antisense |
| 30 | 129  | ATGGAGTACATGGTCTAC-<br>TTCAACTTCTTGTGTGGG-<br>TGCTGCCCGCT-<br>(SEQ ID NO. 14) | ---      | ---       | sense     |

|    | name | sequence  | position     | clone                 | direction |
|----|------|---|--------------|-----------------------|-----------|
| 5  | 130  | GAAGATCCGCAAATAGACA-<br>CCCAGCATGAGCAGAAGCG-<br>GGGGCAGCACCC<br>(SEQ ID NO. 15) | ---          | ---                   | antisense |
| 10 | 131  | CCCTCTAGAGCCCAGCCTGT-<br>GCCCGCCATGCCCATCATGG-<br>GCTCC (SEQ ID NO. 16)         | 2-19<br>1-14 | 5'hva2-29<br>5'hva1-9 | sense     |
| 15 | lamL | CCCACCTTTGAGCAAGTTC<br>(SEQ ID NO. 17)  | ---          | λt10                  | ---       |
| 20 | lamR | GGCTTATGAGTATTCTTCC<br>(SEQ ID NO. 18)  | ---          | λt10                  | ---       |
| 25 | 207  | CCCAAGCTTATGAAAGCCAA<br>CAATACC (SEQ ID NO. 27)                                 | ---          | ---                   |           |
|    | 208  | TGCTCTAGACTCTGGTATCT<br>TCACATT (SEQ ID NO. 28)                                 | ---          | ---                   |           |

### EXAMPLE 2

#### Human A1 adenosine receptor expression construct:

To express the human adenosine receptor cDNA in COS,  
30 CHO and HEK 293 cells, the 118bp SalI-SmaI fragment of the human ventricle A1 PCR product (5'HVA1-9) was ligated together with the 1.8 SmaI-EcoRI fragment of the human kidney A1 adenosine receptor cDNA (hkA1-14) and the 3.0 kb SalI-EcoRI fragment of

pBLUESCRIPT II KS+, resulting in a plasmid containing the contiguous full length coding sequence for the human A1 adenosine receptor cDNA and some 5' and 3' untranslated sequence. This plasmid was digested first with EcoRI, the resulting ends were filled in by Klenow enzyme extension and then the plasmid was digested with XhoI to release a fragment of 1.9 kb containing the full length human A1 adenosine receptor cDNA. The fragment was subcloned into the expression vector pSVL (PHARMACIA) which had been digested with XhoI-SmaI.

Human A2a adenosine receptor expression construct:

To express the human A2a adenosine receptor cDNA in COS, CHO or HEK 293 cells, a contiguous A2a cDNA sequence was constructed before subcloning into the expression vector, pSVL. Primer 131, containing an XbaI recognition site, 14 bp of 5' untranslated sequence of human A1 adenosine receptor cDNA, and the first 18 bp of human A2a adenosine receptor cDNA coding sequence was used with primer 75 in PCR with 1 ng of the plasmid containing the human ventricle A2a 2nd round RACE product (5'hvA2-29) as template. Twenty-five cycles of 40 sec at 94°C, 1 min at 55°C, and 3 min at 72°C, then a final incubation of 15 min at 72°C, with 1 ng of plasmid template and 50 pmol of each primer in a volume of 50 µL according to the GENEAMP protocol (PERKIN ELMER CETUS, Norwalk, CT), resulted in the expected 302 bp product determined by agarose gel electrophoresis. The 172 bp XbaI-EagI digestion product of this DNA fragment was ligated together with 1125 bp EagI-BglII digestion product of the human striata A2a adenosine receptor cDNA (hbA2-22A) and XbaI-SmaI digested pSVL (PHARMACIA), generating the full length human A2a adenosine receptor cDNA coding sequence in a plasmid suitable for its expression in COS, CHO or HEK 293 cells.

Mammalian cell expression:

COS7 cells (ATCC #1651-CRL) were grown in complete medium, Dulbecco's modified Eagle's medium, DMEM (GIBCO, Grand Island, NY) containing 10% fetal bovine serum, 100U/mL

penicillin-streptomycin and 2 mM glutamine, in 5% CO<sub>2</sub> at 37°C. Transient transfection of COS7 cells was performed by the CaPO<sub>4</sub> method (Graham,F.L. and Van Der Erb, A.J. (1973) Virology 52:456-567) using the Mammalian Transfection Kit (STRATAGENE). See Chen and Okayama Mol. Cell Biol. 7:2745-2752. Plasmid DNA (15 mg) was precipitated with 125 mM CaCl<sub>2</sub> in BBS (N,N-bis(2-hydroxyethyl)-2-aminoethanesulfonic acid buffered saline) at room temperature for 30 minutes. The DNA precipitate was added to the COS7 cells and incubated for 18h in 5% CO<sub>2</sub> at 37°C. The precipitate 5 was removed and the cells were washed twice with serum free DMEM. Cells were incubated in complete medium in 5% CO<sub>2</sub> at 37°C for 48 h prior to the binding assay.

Stable expression in CHO or HEK 293 cells:

To establish stable cell lines, CHO or HEK 293 cells were co-transfected with 20 µg of pSVL containing the adenosine receptor cDNA and 1mg of pWLneo (STRATAGENE) containing the neomycin gene. See Southern and Berg (1982) J. Mol. App. Gen. 1:327-341. Transfection was performed by the CaPO<sub>4</sub> method. DNA was precipitated at room temperature for 30 minutes, added to the CHO 15 cells and incubated 18h in 5% CO<sub>2</sub> at 37°C. The precipitate was removed and the cells were washed twice with serum free DMEM. Cells were incubated for 24h in 5% CO<sub>2</sub> at 37°C, replated in 24-well dishes at a dilution of 1:10, and incubated an additional 24h before adding selection medium, DMEM containing 10% fetal bovine serum, 20 100U/mL penicillin-streptomycin, 2 mM glutamine and 0.5 mg/mL G418 (GIBCO). Transfected cells were incubated at 5% CO<sub>2</sub>, 37°C until viable colonies were visible, approximately 14-21 days. Colonies 25 were selected and propagated. The cell clone with the highest number of human adenosine receptors was selected for subsequent application in the binding assay.

EXAMPLE 3

Binding studies:

Membranes were prepared from transiently transfected COS7 cells 48 h after transfection or from G418-selected stably transfected CHO or HEK 293 cells. Cells were harvested in 1 mM EDTA in phosphate buffered saline and centrifuged at 2000 x g for 10 minutes. The cell pellet was washed once with phosphate buffered saline. The cell pellet was resuspended in 2 mL of 5 mM Tris, pH 7.6/5 mM MgCl<sub>2</sub>. Membranes were prepared from the cells by freeze-thaw lysis in which the suspension was frozen in a dry ice/ethanol bath and thawed at 25°C twice. The suspension was homogenized after adding an additional 2 mL of 5 mM Tris, pH 7.6/5 mM MgCl<sub>2</sub>, in a glass dounce homogenizer with 20 strokes. The membranes were pelleted at 40,000 x g at 4°C for 20 minutes. The membrane pellet was resuspended at a protein concentration of 1-2 mg/mL in binding assay buffer, 50 mM Tris, pH 7.6/10 mM MgCl<sub>2</sub>. Protein concentration was determined by the method of Bradford ((1976) Anal. Biochem. 72: 248-250). Before the binding assay was performed, the membranes were incubated with adenosine deaminase (BOEHRINGER MANNHEIM), 2 U/mL for 30 minutes at 37°C. Saturation binding of [<sup>3</sup>H]-cyclohexyladenosine (CHA) was performed on membranes prepared from pSVLA1 transfected COS7 or CHO cells.

Membranes (100µg) were incubated in the presence of 0.2 U/mL adenosine deaminase with increasing concentrations of CHA (NEN, 32 Ci/mmol) in the range of 0.62 - 30 nM for 120 minutes at 25°C in a total volume of 500 µL. The binding assay was terminated by rapid filtration and three washes with ice-cold 50 mM Tris, pH 7.6/10 mM MgCl<sub>2</sub> on a SKATRON CELL HARVESTER equipped with a receptor binding filtermat (SKATRON INSTRUMENTS, INC). Non-specific binding was determined in the presence of 100 µM N<sup>6</sup>-cyclopentyladenosine (CPA). Bound radioactivity was measured by scintillation counting in READY SAFE SCINTILLATION COCKTAIL (BECKMAN). For competition binding experiments, membranes were

incubated with 5 nM [<sup>3</sup>H]-CHA and various concentrations of A1 adenosine receptor agonists. Saturation binding of [<sup>3</sup>H] CGS-21680 was performed on membranes prepared from pSVLA2a transfected COS7 cells. Membranes (100 $\mu$ g) were incubated in the presence of 0.2 U/mL adenosine deaminase with increasing concentrations of CGS21680 (NEN, 48.6 Ci/mmol) in the range of 0.62 -80 nM for 90 minutes at 25°C in a total volume of 500  $\mu$ L. The binding assay was terminated by rapid filtration with three washes with ice-cold 50 mM Tris, pH 7.6/10 mM MgCl<sub>2</sub> on a Skatron cell harvester equipped with a receptor binding filtermat (SKATRON INSTRUMENTS, INC). Non-specific binding was determined in the presence of 100 $\mu$ M CPA. Bound radioactivity was measured by scintillation counting in READY SAFE LIQUID SCINTILLATION COCKTAIL (BECKMAN). For competition binding experiments, membranes were incubated with 5nM [<sup>3</sup>H]-CGS21680 and various concentrations of A2 adenosine receptor agonists.

Saturation binding of [<sup>3</sup>H]5'-N-ethylcarboxamidoadenosine (NECA) was performed on membranes (100  $\mu$ g) prepared from pSVLhb32C (A2b) transfected COS7 cells in the presence of adenosine deaminase with increasing concentrations of NECA (NEN, 15.1Ci/mmol) in the range of 1.3-106 nM for 90 minutes at 25°C in a total volume of 500  $\mu$ L. The assay was terminated by rapid filtration and three washes with ice-cold binding buffer on a cell harvester equipped with a receptor binding filtermat (SKATRON INSTRUMENTS, INC). Bound radioactivity was measured by scintillation counting. Non-specific binding was measured on membranes prepared from non-transfected COS7 cells. For competition binding experiments, membranes from transfected cells were incubated with 10 nM [<sup>3</sup>H]NECA and varying concentrations of adenosine receptor antagonists.

EXAMPLE 4

The human A3 adenosine receptor was cloned from a human striata cDNA library. Oligonucleotide probes were designed based on the rat A3 sequence of Zhou et al., Proc. Natl. Acad. Sci. 89, 7432 (1992). The complete sequence of the human A3 adenosine receptor was determined and the protein sequence deduced. The cloned human A3 adenosine receptor is expressed in a heterologous expression system in COS, CHO and HEK 293 cells. Radiolabeled adenosine receptor agonists and antagonists are used to measure the binding properties of the expressed receptor. Stable cell lines can be used to evaluate and identify adenosine receptor agonists, antagonists and enhancers.

STEP A:

A synthetic probe homologous to the rat A3 adenosine receptor was generated using the polymerase chain reaction (PCR). Three  $\mu$ l of rat brain cDNA was used as template in a PCR amplification reaction according to the GENEAMP protocol (PERKIN ELMER CETUS, Norwalk, CT) containing 50 pmol of primers 207 (5'-cccaagcttatgaaagccaacaatacc) (SEQ. ID NO: 27) and 208 (5'-tgctctagactctggtatcttcacatt) (SEQ. ID NO: 28) in a total volume of 50 ml. Primers 207 and 208 are based on the published rat A3 adenosine receptor sequence (Zhou, et al, (1992), Proc. Natl. Acad. Sci. USA, 89:7432-7406). Forty cycles of 40 sec at 94°C, 1 min at 55°C, 3 min at 72°C were performed and the resulting 788 bp fragment was subcloned into HindIII-XbaI digested pBLUESCRIPT II KS+ (STRATAGENE, La Jolla, CA). The sequence was verified by the SEQUENASE protocol (USBC, Cleveland, OH).

STEP B:

The 788 bp PCR fragment was labeled with  $\alpha^{32}P$ -dCTP using the MULTIPRIME DNA LABELLING SYSTEM (AMERSHAM, Arlington Heights, IL) and used to screen a human striata cDNA library

(STRATAGENE, La Jolla, CA). E. coli strain XL-1 Blue (STRATAGENE, La Jolla, CA) cells were infected with library phage and grown overnight at 37°C. Phage DNA was transferred to HYBOND-N nylon filters according to the manufacturer's protocol (AMERSHAM, Arlington Heights, IL). The probe was incubated with the filters in 5 X SSC, 30% formamide, 5 X Denhardt's solution, 0.5% sodium dodecyl sulfate, and 50 mg/ml sonicated salmon testis DNA. The filters were washed in 2 X SSC at 55°C. A positively hybridizing phage (HS-21a) was identified and plaque purified by two additional rounds of plating and hybridization. The insert was subcloned to the plasmid pBLUESCRIPT II SK- according to the manufacturer's protocol (STRATAGENE, La Jolla, CA). Upon sequence analysis using the SEQUENASE protocol (USBC, Cleveland, OH) it was determined that clone HS-21a contained the complete open reading frame corresponding to the human homolog of the rat A3 adenosine receptor. The coding region of the human A3 adenosine receptor cDNA is 78% identical to the rat sequence at the nucleotide level and contains 265 bp and 517 bp of 5' and 3' untranslated sequence, respectively. The 1.7 kb fragment was excised using sites present in the multiple cloning site of pBLUESCRIPT II SK- (STRATAGENE, La Jolla, CA) and subcloned into Xhol/SacI digested pSVL (PHARMACIA, Piscataway, NJ) for its expression in COS and CHO cells.

#### EXAMPLE 5

25 Mammalian cell expression:

COS7 cells (ATCC #1651-CRL) were grown in complete medium, Dulbecco's modified Eagle's medium, DMEM (GIBCO, Grand Island, NY) containing 10% fetal bovine serum, 100U/mL penicillin-streptomycin and 2mM glutamine, in 5% CO<sub>2</sub> at 37°C. Transient transfection of COS7 cells was performed by the CaPO<sub>4</sub> method (Graham, F.L. and Van Der Eb, A.J. (1973) Virology 52:456-567) using the Mammalian Transfection Kit (STRATAGENE). Plasmid DNA (15 mg) was precipitated with 125 mM CaCl<sub>2</sub> in BBS (N,N-bis(2-

hydroxyethyl)-2-aminoethanesulfonic acid buffered saline) at room temperature for 30 minutes. The DNA precipitate was added to the COS7 cells and incubated for 18 h in 5% CO<sub>2</sub> at 37°C. The precipitate was removed and the cells were washed twice with serum free DMEM. Cells were incubated in complete medium in 5% CO<sub>2</sub> at 37°C for 48 h prior to the binding assay.

Stable expression in CHO cells:

To establish stable cell lines, CHO cells were cotransfected with 20 µg of pSVL containing the adenosine receptor cDNA and 1 µg of pWLneo (STRATAGENE) containing the neomycin gene. Transfection was performed by the CaPO<sub>4</sub> method. DNA was precipitated at room temperature for 30 minutes, added to the COS7 cells and incubated 18 h in 5% CO<sub>2</sub> at 37°C. The precipitate was removed and the cells were washed twice with serum free DMEM. Cells were incubated for 24 h in 5% CO<sub>2</sub> at 37°C, replated in 24-well dishes at a dilution of 1:10, and incubated an additional 24 h before adding selection medium, DMEM containing 10% fetal bovine serum, 100U/mL penicillin-streptomycin, 2 mM glutamine and 1.0 mg/mL G418 (GIBCO). Transfected cells were incubated at 5% CO<sub>2</sub>, 37°C until viable colonies were visible, approximately 14-21 days. Colonies were selected and propagated. The cell clone with the highest number of human adenosine receptors was selected for subsequent application in the binding assay.

25

EXAMPLE 6

Binding assay:

Membranes were prepared from transiently transfected COS7 cells 48 h after transfection or from G418-selected stably transfected CHO or HEK 293 cells. Cells were harvested in 1 mM EDTA in phosphate buffered saline and centrifuged at 2000 x g for 10 minutes. The cell pellet was washed once with phosphate buffered saline. The cell pellet was resuspended in 2 mL of 5 mM Tris, pH 7.6/

5 5mM MgCl<sub>2</sub>. Membranes were prepared from the cells by freeze-thaw lysis in which the suspension was frozen in a dry ice/ethanol bath and thawed at 25°C twice. The suspension was homogenized after adding an additional 2 mL of 5 mM Tris, pH 7.6/ 5mM MgCl<sub>2</sub>, in a glass dounce homogenizer with 20 strokes. The membranes were pelleted at 40,000  
10 x g at 4°C for 20 minutes. The membrane pellet was resuspended at a protein concentration of 1-2 mg/mL in binding assay buffer, 50 mM Tris, pH 7.6/10 mM MgCl<sub>2</sub>. Protein concentration was determined by the method of Bradford ((1976) Anal. Biochem. 72: 248-250). Before the binding assay was performed, the membranes were incubated with  
adenosine deaminase (BOEHRINGER MANNHEIM), 2U/mL for 30  
15 minutes at 37°C. Saturation binding of [<sup>125</sup>I]-N<sup>6</sup>-aminobenzyl-adenosine (<sup>125</sup>I-ABA) or [<sup>125</sup>I]-N<sup>6</sup>-2-(4-amino-3-iodophenyl)ethyl-adenosine (APNEA) was performed on membranes prepared from pSVLA3 transfected COS7 cells. Membranes (100 µg) were incubated  
in the presence of 0.2U/mL adenosine deaminase with increasing concentrations of <sup>125</sup>I-ABA in the range of 0.1-30 nM for 120 minutes  
20 at 25°C in a total volume of 500 µL. The binding assay was terminated by rapid filtration and three washes with ice-cold 50 mM Tris, pH 7.6/10 mM MgCl<sub>2</sub> on a Skatron cell harvester equipped with a receptor binding filtermat (SKATRON INSTRUMENTS, INC). Non-specific binding was determined on non-transfected cells. Bound radioactivity was measured by scintillation counting in Ready Safe Scintillation Cocktail (BECKMAN).

25

#### EXAMPLE 7

30 In vitro transcription and oocyte expression:  
The 1.3 kb XhoI-BamHI fragment of the pSVL expression construct (described in Example 2) containing the full length human A2a adenosine receptor coding sequence was ligated into SalI-SpeI digested pGEMA (Swanson, et al, (1990) Neuron 4:929-939). The resulting plasmid, pGEMA2, was linearized with NotI, forming a template for in vitro transcription with T7 RNA polymerase. The

homologous adenosine receptor subtype cDNA in pBluescript SK- was used as a template for in vitro transcription by T3 polymerase after removal of most of the 5' untranslated region, with the exception of 20 bp, as a 0.3 kb SmaI fragment. The K<sup>+</sup> channel cDNA, Kv3.2b was employed as a negative control in the cAMP accumulation assay. The generation of Kv3.2b RNA was described by Luneau, et al, ((1991) FEBS Letters 1:163-167). Linearized plasmid templates were used with the STRATAGENE mCAP kit according to the manufacturer's protocol, except that the SP6 RNA polymerase reaction was performed at 40°C. Oocytes were harvested from mature female *Xenopus laevis*, treated with collagenase, and maintained at 18°C in ND96 medium (GIBCO) supplemented with 1 mM sodium pyruvate and 100 mg/mL gentamycin. Fifty nanoliters (10 ng) of RNA diluted in H<sub>2</sub>O was injected and oocytes were incubated at 18°C for 48 hours.

15

#### EXAMPLE 8

##### cAMP accumulation assay in oocytes:

Oocytes injected with either human adenosine receptor transcript or the Kv3.2b transcript were transferred to fresh medium supplemented with 1 mM of the phosphodiesterase inhibitor, Ro 20-1724 (RBI, Natick, MA) and 1 mg/mL bovine serum albumin incubated for 30 minutes and transferred to an identical medium with or without the agonist adenosine (10 mM) for an additional 30 minutes at room temperature. Groups of 5-10 oocytes were lysed by transfer to ND96/100 mM HCl/1 mM Ro 20-1724 in microfuge tubes, shaken, incubated at 95°C for 3 min, and centrifuged at 12000 g for 5 min. Supernatants were stored at -70°C before cAMP measurements. Cyclic AMP levels were determined by radioimmunoassay (RIANEN kit, DuPont/NEN) using the acetylation protocol. The adenosine receptor antagonist, 8-(p-sulfophenyl)theophylline (100 μM) was utilized to inhibit the cAMP response induced by adenosine in oocytes expressing the adenosine receptors.

EXAMPLE 9

cAMP accumulation in stable CHO cell lines:

The changes in cAMP accumulation can alternatively be measured in stably transfected CHO cells expressing the human adenosine receptor subtypes. CHO cells are washed twice in phosphate buffered saline (PBS) and detached in 0.2% EDTA in PBS. The cells are pelleted at 800 rpm for 10 min and resuspended in KRH buffer (140 mM NaCl/5 mM KCl/2 mM CaCl<sub>2</sub>/1.2 mM MgSO<sub>4</sub>/1.2 mM KH<sub>2</sub>PO<sub>4</sub>/6 mM glucose/25 mM Hepes buffer, pH 7.4). The cells are washed once in KRH buffer and resuspended at 10<sup>7</sup> cells/mL. The cell suspension (100 µL) is mixed with 100 µL of KRH buffer containing 200 mM Ro 20-1724 and incubated at 37°C for 10 minutes. Adenosine (10 mM), NECA or CPCa was added in 200 µL KRH buffer containing 200 µM Ro 20-1724 and incubated at 37°C for 20 minutes. After the incubation, 400 mL of 0.5 mM NaOAc (pH 6.2) was added and the sample was boiled for 20 minutes. The supernatant was recovered by centrifugation for 15 minutes and stored at -70°C. cAMP levels were determined by radioimmunoassay (RIANEN kit, DuPont/NEN) using the acetylation protocol. The effect of antagonists on cAMP accumulation are measured by preincubation for 20 minutes before adding adenosine.

EXAMPLE 10

Expression Construct and Transfection

The 1.7 kb HS-21a cDNA (A3) was subcloned as a Sal-I-BamHI fragment into the expression vector pCMV5 (Mumby, S.M., Heukeroth, R.O., Gordon, J.I. and Gilman, A.G. (1990) Proc. Natl. Acad. Sci. USA 87, 728-732.) creating the vector pCMV5-A3. CHO or HEK 293 cells stably expressing the human HS-21a cDNA were prepared by co-transfection of 15 µg pCMV5-A3 and 1 µg pWLneo (Stratagene) using the calcium phosphate method. Stable cell lines were also generated using EBV based mammalian expression vectors, pREP (INVITROGEN). Neomycin resistant colonies were selected in 1

mg/mL G418 (GIBCO). Stable colonies were screened for expression of HS-21a by  $^{125}\text{I}$ -ABA binding.

EXAMPLE 11

Binding Studies

Membranes were prepared from stable CHO cell lines in 10 mM Hepes, pH 7.4 containing 0.1 mM benzamidine and 0.1 mM PMSF as described (Mahan, L.C., et al., (1991) Mol. Pharmacol. 40, 1-7). Pellets were resuspended in 5 mM Hepes, pH 7.4/5 mM MgCl<sub>2</sub>/0.1 mM benzamidine/0.1 mM PMSF at a protein concentration of 1-2 mg/mL and were incubated with adenosine deaminase (Boehringer Mannheim), 2U/mL at 37 °C for 20 minutes. Saturation binding of  $^{125}\text{I}$ -ABA was carried out on 50 mg of membranes for 120 minutes at 25 °C in a total volume of 100  $\mu\text{L}$ . The assay was terminated by rapid filtration and three washes with ice-cold binding buffer on a Skatron harvester equipped with a receptor binding filtermat (Skatron Instruments, INC). The specific activity of  $^{125}\text{I}$ -ABA, initially 2,200 Ci/mmol, was reduced to 100 Ci/mmol with nonradioactive I-ABA for saturation analysis. Nonspecific binding was measured in the presence of 1 mM I-ABA. The K<sub>D</sub> and B<sub>max</sub> values were calculated by the EBDA program (McPherson, G.A. (1983) Computer Programs for Biomedicine 17, 107-114). Competition binding of agonists and antagonists was determined with  $^{125}\text{I}$ -ABA (0.17-2.0 nM, 2000 Ci/mmol). Nonspecific binding was measured in the presence of 400 mM NECA. Binding data were analyzed and competition curves were constructed by use of the nonlinear regression curve fitting program Graph PAD InPlot, Version 3.0 (Graph Pad Software, San Diego). K<sub>i</sub> values were calculated using the Cheng-Prusoff derivation (Cheng, Y.C. and Prusoff, H.R. (1973) Biochem. Pharmacol. 22, 3099-3108.).

The binding properties of the receptor encoded by HS-21a were evaluated on membranes prepared from CHO cells stably expressing the HS-21a cDNA. The radioligand,  $^{125}\text{I}$ -APNEA, was previously used to characterize rat A<sub>3</sub> adenosine receptors. In preliminary experiments, high non-specific  $^{125}\text{I}$ -APNEA binding to

CHO cell membranes was observed which interfered with the measurement of specific binding to expressed receptors. Specific and saturable binding of the adenosine receptor agonist,  $^{125}\text{I}$ -ABA was measured on membranes prepared from the stably transfected cells (Figure 11A). The specific binding of  $^{125}\text{I}$ -ABA could be prevented by either 1 mM nonradioactive I-ABA or 400  $\mu\text{M}$  NECA. No specific binding of  $^{125}\text{I}$ -ABA was measured on membranes prepared from non-transfected CHO cells. The specific binding of  $^{125}\text{I}$ -ABA measured in either the presence of 10  $\mu\text{M}$  GTP $\gamma$ S or 100  $\mu\text{M}$  Gpp(NH)p was reduced by 56 and 44% respectively, relative to the specific binding measured in the absence of the uncoupling reagents. These results suggest that  $^{125}\text{I}$ -ABA exhibits some agonist activity on the receptor encoded by the HS-21a cDNA expressed in the stable CHO cell line.  $^{125}\text{I}$ -ABA binds to membranes prepared from the HS-21a stable CHO cells with a dissociation constant of 10 nM ( $B_{\max} = 258 \text{ fmol/mg protein}$ ) with a Hill coefficient of 0.99 indicating binding to a single class of high affinity sites (Figure 11B).

The competition of adenosine receptor agonists and antagonists for binding to HS-21a receptors was determined (Figure 12). The  $K_i$  values for agonists (top panel) were calculated to be 26 nM for NECA, 34 nM for R-PIA, 89 nM for CPA and 320 nM for S-PIA, resulting in a potency order profile of NECA > R-PIA > CPA > S-PIA. In contrast to the insensitivity of adenosine receptor antagonists reported for the rat A3 adenosine receptor subtype, a number of xanthine antagonists exhibited competition with  $^{125}\text{I}$ -ABA for binding to the HS-21a receptor (Figure 12, lower panel). Studies of the sheep A3 adenosine receptor indicated that 8-phenylxanthines substituted in the para-position with acidic substituents are high affinity antagonists. By evaluating additional xanthines in this class I-ABOPX was determined to be the highest affinity antagonist yet reported for A3 adenosine receptors. The  $K_i$  values for antagonists were calculated to be 18 nM for I-ABOPX, 55 nM for BW-A1433, 70 nM for XAC and 750 nM for DPCPX, resulting in a potency order profile of I-ABOPX > BW-A1433 > XAC > DPCPX.

EXAMPLE 12

cAMP Studies

Determinations were made on stably transfected CHO cells in suspension as described (Linden et al., (1993) Mol. Pharm. 44:524-532). Supernatants (500 µL) were acetylated and acetylcylic AMP was measured by automated radioimmunoassay (Hamilton, B.R. and Smith, D.O. (1991) J. Physiol. (Lond.) 432, 327-341). Antagonist dissociation constants were estimated from pA<sub>2</sub> values as described by Schild (1957) Pharm. Rev. 9, 242-246).

10

EXAMPLE 13

Northern Blot Analysis

Human poly(A)<sup>+</sup> RNA from different tissue sources (Clontech) is fractionated on a 1% agarose-formaldehyde gel (Sambrook, J., Fritsch, E. and Maniatis, T. (1989) *Molecular Cloning: A Laboratory Manual*, Second Edition, (Cold Spring Harbor Press, Cold Spring Harbor, NY), transferred to Hybond-N membranes and hybridized in 5XSSPE, 5XDenhardt's, 0.5% SDS, 50 mg/mL sonicated salmon testis DNA, with 30% formamide (for A1, A2a, and A2b) or 50% formamide (for HS-21a) at 42°C. DNA probes corresponding to nucleotides 512-1614, 936-2168, and 321-1540 of accession numbers X68485(A1), X68486(A2a), and X68487(A2b) respectively, and a 1.7 kb SalI-BamHI fragment of HS-21a were labeled with  $\alpha^{32}P$ -dCTP by the random priming method. Filters were washed under high stringency conditions in 0.1XSSC at 65°C.

EXAMPLE 14

INHIBITION OF TNF $\alpha$  PRODUCTION

30      STEP A:

Isolation of human peripheral blood mononuclear cells.

Human blood is obtained by venipuncture from healthy donors and collected into tubes containing 20U/mL of heparin sodium salt. The blood is diluted 1:1 with Hanks balanced salts solution containing 20 U/mL Heparin. Peripheral blood mononuclear cells (PBMC) are isolated by Ficoll-Hypaque density centrifugation. The PBMC are resuspended in a small volume (2-5 mL) of RPMI + 10% autologous human serum, counted then diluted further with RPMI + 10% autologous human serum to 5 x 10<sup>5</sup> cells/mL. Subsequently the cells are plated in a six well Costar plastic plate precoated with 1 mg / mL fibronectin. Lipopolysaccharide, as well as the appropriate adenosine agonists and antagonists, are added simultaneously. Following incubation at 37°C for 18 hours, the cell culture supernatants are harvested, clarified and tested for TNF levels by a specific trapping ELISA.

15 STEP B:  
ELISA for human TNF $\alpha$ .

A mouse anti-human TNF $\alpha$  monoclonal antibody is diluted to 0.5 mg/mL in PBS - MgCl<sub>2</sub> - CaCl<sub>2</sub> and added to plastic 96 - well plates. Following a 24 hr incubation at 4°C the plates are washed with PBS-Tween then treated with a solution of PBS and 1% BSA. Following additional washing with PBS Tween, aliquots of monocytes thought to contain TNF $\alpha$  are added to the dish, diluted to 100 mL with PBS tween and incubated for 2 hours at 37°C. The plates are further washed with PBS-Tween, then treated with a 1 to 2000 dilution of rabbit anti-human TNF polyclonal antiserum (Genzyme). The plates are incubated for 1 hour, washed then treated again with a goat anti-rabbit IgG Fab-horseradish peroxidase conjugate. The plates are incubated for one hour, washed, and the bound peroxidase is detected by the addition of a TMB peroxide mixture. TNF $\alpha$  levels are determined by comparison with a standard curve generated using pure recombinant TNF $\alpha$ .

EXAMPLE 15

DETECTION OF ADENOSINE RECEPTOR TRANSCRIPTS BY  
REVERSE-TRANSCRIPTASE POLYMERASE CHAIN REACTION  
AMPLIFICATION

STEP A:

5        Total RNA was extracted by the guanidinium isothiocyanate method (Chirgwin, J.M., et al, (1979) Biochemistry 18:5294-5299) from normal and LPS-stimulated human monocytes. First strand cDNA was reverse transcribed from 600 ng total RNA in a volume of 20 ml containing 20mM Tris-HCL (pH 8.4), 50mM KCl, 2.5mM MgCl<sub>2</sub>, 0.1mg/ml bovine serum albumin (BSA), 0.5mM dNTP's, 10 mM DTT,  
10      10 units SUPERSCRIPT II reverse transcriptase (LIFE TECHNOLOGIES, INC., Gathersburg, MD), and 50ng random hexamers.

STEP B:

15      Human adenosine receptor subtype transcript expression was determined using the polymerase chain reaction (PCR). Three µl of the randomly primed first strand cDNA, prepared from monocytes (+) or (-) LPS stimulation, was used as template in a PCR amplification reaction according to the GENEAMP protocol (PERKIN ELMER CETUS, Norwalk, CT) containing 50pmol subtype selective primers in a total volume of 100 µl. Primer pairs were designed to span four (A1 primers) and five (A2a, A2b, A3 primers) transmembrane domains and gave no or incorrect sized PCR products when tested on human genomic DNA. Primer pairs for amplification (see Table 1) were 266+267 (A1),  
20      253+254 (A2a), 261+262 (A2b), 230+236 (A3), and 141+142 for glyceraldehyde-3-phosphate dehydrogenase (GAPDH). Primers 141+142 are based on the published human GAPDH sequence (Tokunaga, K., et al, (1987) Cancer Research 47:5616-5619). Cycling parameters were 1 min at 94°C, 1 min at 55°C, 3 min at 72°C for 35 cycles (A1), 25 cycles (A2a), 35 cycles (A3), and 20 cycles (GAPDH).  
25      Cycling parameters for A2b were 1 min at 94°C, 1 min at 59°C, 3 min at 72°C for 30 cycles.  
30

STEP C:

Ten  $\mu$ l of each PCR amplification reaction was electrophoresed on a 1.4% agarose gel and alkaline blotted to Zeta-Probe GT membranes according to the manufacturer's protocol (BIO-RAD, Hercules, CA). Membranes were hybridized in 0.25 M sodium phosphate (pH 7.2), 0.5M NaCl, 7.0% sodium dodecyl sulphate (SDS), 1 mM EDTA, 1% BSA, and  $1 \times 10^6$  cpm/ml  $^{32}$ P labeled probe at 50°C. Double-stranded DNA probes were generated by Klenow enzyme extension of annealed oligonucleotide pairs including a $^{32}$ P-dCTP.

Oligonucleotide pairs for probe synthesis (see Table 1) were 268+269 (A1), 66+67 (A2a), 263+264 (A2b), 259+260 (A3), and 143+144 (GAPDH). Oligonucleotides 259+260 are based on the published sheep A3 adenosine receptor (Linden, J., et al, (1993) Molecular Pharmacology 44:524-532) and 143+144 on the human GAPDH sequence (Tokunaga et al). Following hybridization membranes were washed to a final stringency of 75mM NaCl, 7.5mM sodium citrate, 0.1% SDS and exposed to autoradiography film. All four adenosine receptor subtypes were found to be present on monocytes through this analysis.

20

25

30

TABLE 1:

| <u>NAME</u> | <u>SEQUENCE</u>   |
|-------------|---|
| 66          | 5' GCCTCTTGAGGATGTGGTCCCCATGAACACTACATGGTGTACTTCA               |
| 5 67        | 5' GCAGGGGCACCAGCACACAGGCAAAGAAGTTGAAGTACACCATGT                |
| 141         | 5' TCACCATCTTCCAGGAGC   |
| 142         | 5' ACTCCTTGGAGGCCATGT   |
| 143         | 5' TCCTGCACCACCAACTGCTTAGCCCCCTGGCCAAGGTATCCAT                  |
| 10 144      | 5' CATGAGCCCTTCCACGATGCCAAAGTTGTATGGATGACCTTGGC                 |
| 230         | 5' GTTACCTACATCACCATG   |
| 236         | 5' GTTAGATAAGTTCAGACT   |
| 253         | 5' TCCTCGGTGTACATCACG   |
| 15 254      | 5' TCCATCTGCTTCAGCTGT   |
| 259         | 5' CTGGGCCTTGCTGGCTGGTGTCAATTCTGGTGGATTGACCCCC                  |
| 260         | 5' TGAGGTCAAGTTCATGTTCCAGCCAAACATGGGGTCAATCCCAC                 |
| 261         | 5' ATGCTGCTGGAGACACAGGA   |
| 20 262      | 5' TGGTCCATCAGCTCAGTGC  |
| 263         | 5'<br>GGTGGAACAGTAAAGACAGTGCCACCAACAATGCACAGAACCTGGATGGAACCACGA |
| 264         | 5' GGACCACATTCTCAAAGAGACACTTCACAAGGCAGCAGCTTCATTGTGGTCCATCCC    |
| 266         | 5' CTACATCGGCATCGAGGT   |
| 25 267      | 5' GAACTCGCACITGATCAC   |
| 268         | 5' TGGTGGACTGACCCCTATGTTGGCTGGAACAATCTGAGTGC GG                 |
| 269         | 5' TGCTGCCGTTGGCTGCCAGGCCGCTCCACCGCACTCAGATTGT                  |

30 While the foregoing specification teaches the principles of the present invention, with examples provided for the purpose of illustration, it will be understood that the practice of the invention encompasses all of the usual variations, adaptations, modifications, as come within the scope of the following claims and its equivalents.

SEQUENCE LISTING

(1) GENERAL INFORMATION:

(i) APPLICANT: Jacobson, Marlene A

5 (ii) TITLE OF INVENTION: INHIBITION OF TNFalpha PRODUCTION  
BY A2b ADENOSINE RECEPTOR AGONISTS AND ENHANCERS

(iii) NUMBER OF SEQUENCES: 56

(iv) CORRESPONDENCE ADDRESS:

- (A) ADDRESSEE: Merck & Co., Inc.
- (B) STREET: P.O.Box 2000
- (C) CITY: Rahway
- 10 (D) STATE: New Jersey
- (E) COUNTRY: United States
- (F) ZIP: 07065

(v) COMPUTER READABLE FORM:

- (A) MEDIUM TYPE: Floppy disk
- (B) COMPUTER: IBM PC compatible
- (C) OPERATING SYSTEM: PC-DOS/MS-DOS
- 15 (D) SOFTWARE: PatentIn Release #1.0, Version #1.25

(vi) CURRENT APPLICATION DATA:

- (A) APPLICATION NUMBER:
- (B) FILING DATE: 6-MAY-1994
- (C) CLASSIFICATION:

(viii) ATTORNEY/AGENT INFORMATION:

- 20 (A) NAME: Bencen, Gerard H  
(B) REGISTRATION NUMBER: 35,746  
(C) REFERENCE/DOCKET NUMBER: 19222

(ix) TELECOMMUNICATION INFORMATION:

- (A) TELEPHONE: (908) 594-3901
- (B) TELEFAX: (908)594-4720

25 (2) INFORMATION FOR SEQ ID NO:1:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 60 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

30 (ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

(iv) ANTI-SENSE: NO

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

ATTCGCAGCC ACGTCCTGAG GCGGCCGGAG CCCTCAAAG CAGGTGGCAC CAGTGCCCCC

60

(2) INFORMATION FOR SEQ ID NO:2:

5

- (i) SEQUENCE CHARACTERISTICS:  
(A) LENGTH: 60 base pairs  
(B) TYPE: nucleic acid  
(C) STRANDEDNESS: single  
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

10

(iii) HYPOTHETICAL: NO

(iv) ANTI-SENSE: NO

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:

15

GCGGAGGCTG ATCTGCTCTC CATCACTGCC ATGAGCTGCC AAGGCCGGG CACTGGTGCC

60

(2) INFORMATION FOR SEQ ID NO:3:

20

- (i) SEQUENCE CHARACTERISTICS:  
(A) LENGTH: 60 base pairs  
(B) TYPE: nucleic acid  
(C) STRANDEDNESS: single  
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

(iv) ANTI-SENSE: NO

25

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:

TCCAGAAGTT CCGGGTCACC TTCCCTTAAGA TCTGGAATGA CCACCTCCGC TGCCAGCCCA

60

30

(2) INFORMATION FOR SEQ ID NO:4:

- (i) SEQUENCE CHARACTERISTICS:  
(A) LENGTH: 60 base pairs  
(B) TYPE: nucleic acid  
(C) STRANDEDNESS: single  
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

(iv) ANTI-SENSE: NO

5 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:

AGTCGTGGGG CGCCTCCTCT GGGGGTCCCT CGTCGACGGG GGGCGTGGGC TGGCAGCGGA

60

(2) INFORMATION FOR SEQ ID NO:5:

10 (i) SEQUENCE CHARACTERISTICS:  
(A) LENGTH: 45 base pairs  
(B) TYPE: nucleic acid  
(C) STRANDEDNESS: single  
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

15 (iii) HYPOTHETICAL: NO

(iv) ANTI-SENSE: NO

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:

GCCTCTTTGA GGATGTGGTC CCCATGAACCT ACATGGTGTA CTTCA

45

20 (2) INFORMATION FOR SEQ ID NO:6:

(i) SEQUENCE CHARACTERISTICS:  
(A) LENGTH: 45 base pairs  
(B) TYPE: nucleic acid  
(C) STRANDEDNESS: single  
(D) TOPOLOGY: linear

25 (ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

(iv) ANTI-SENSE: NO

30 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:6:

GCAGGGGCAC CAGCACACAG GCAAAGAAAGT TGAAGTACAC CATGT

45

(2) INFORMATION FOR SEQ ID NO:7:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 18 base pairs  
(B) TYPE: nucleic acid  
(C) STRANDEDNESS: single  
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

5 (iii) HYPOTHETICAL: NO

(iv) ANTI-SENSE: NO

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:7:

10 TCGCGCCGCC AGGAAGAT

18

(2) INFORMATION FOR SEQ ID NO:8:

- (i) SEQUENCE CHARACTERISTICS:  
(A) LENGTH: 30 base pairs  
(B) TYPE: nucleic acid  
(C) STRANDEDNESS: single  
(D) TOPOLOGY: linear

15 (ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

(iv) ANTI-SENSE: NO

20 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:8:

TATATTGAAT TCTAGACACC CAGCATGAGC

30

(2) INFORMATION FOR SEQ ID NO:9:

- (i) SEQUENCE CHARACTERISTICS:  
(A) LENGTH: 18 base pairs  
(B) TYPE: nucleic acid  
(C) STRANDEDNESS: single  
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

30 (iv) ANTI-SENSE: NO

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:9:

TCAATGGCGA TGGCCAGG

18

(2) INFORMATION FOR SEQ ID NO:10:

- (i) SEQUENCE CHARACTERISTICS:

  - (A) LENGTH: 30 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear

5

- (ii) MOLECULE TYPE: cDNA
  - (iii) HYPOTHETICAL: NO
  - (iv) ANTI-SENSE: NO

10

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:10:

TATATTGAAT TCATGGAGCT CTGCGTGAGG

30

(2) INFORMATION FOR SEQ ID NO:11:

- 15                   (i) SEQUENCE CHARACTERISTICS:  
                      (A) LENGTH: 18 base pairs  
                      (B) TYPE: nucleic acid  
                      (C) STRANDEDNESS: single  
                      (D) TOPOLOGY: linear

- (ii) MOLECULE TYPE: cDNA
  - (iii) HYPOTHETICAL: NO
  - (iv) ANTI-SENSE: NO

20

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:11:

GTAGACCATG TACTCCAT

18

25

(3) INFORMATION FOR SEQ ID NO:13:

- (i) SEQUENCE CHARACTERISTICS:

  - (A) LENGTH: 30 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear

30

- (ii) MOLECULE TYPE: cDNA  
(iii) HYPOTHETICAL: NO  
(iv) ANTI-SENSE: NO

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:12:

TATATTGAAT TCTGACCTTC TCGAACTCGC

30

(2) INFORMATION FOR SEQ ID NO:13:

- 5 (i) SEQUENCE CHARACTERISTICS:  
(A) LENGTH: 29 base pairs  
(B) TYPE: nucleic acid  
(C) STRANDEDNESS: single  
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

10 (iii) HYPOTHETICAL: NO

(iv) ANTI-SENSE: NO

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:

ATTGAATTCCG ATCACGGGCT CCCCCATGC

29

15 (2) INFORMATION FOR SEQ ID NO:14:

- (i) SEQUENCE CHARACTERISTICS:  
(A) LENGTH: 50 base pairs  
(B) TYPE: nucleic acid  
(C) STRANDEDNESS: single  
(D) TOPOLOGY: linear

20 (ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

(iv) ANTI-SENSE: NO

25 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:14:

ATGGAGTACA TGGTCTACTT CAACTTCTTT GTGTGGGTGC TGCCCCCGCT

50

(2) INFORMATION FOR SEQ ID NO:15:

- 30 (i) SEQUENCE CHARACTERISTICS:  
(A) LENGTH: 50 base pairs  
(B) TYPE: nucleic acid  
(C) STRANDEDNESS: single  
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

(iv) ANTI-SENSE: NO

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:15:

5 GAAGATCCGC AAATAGACAC CCAGCATGAG CAGAAGCGGG GGCAGCACCC

50

(2) INFORMATION FOR SEQ ID NO:16:

- (i) SEQUENCE CHARACTERISTICS:  
(A) LENGTH: 45 base pairs  
(B) TYPE: nucleic acid  
(C) STRANDEDNESS: single  
(D) TOPOLOGY: linear

10 (ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

(iv) ANTI-SENSE: NO

15 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:16:

CCCTCTAGAG CCCAGCCTGT GCCCGCCATG CCCATCATGG GCTCC

45

(2) INFORMATION FOR SEQ ID NO:17:

- (i) SEQUENCE CHARACTERISTICS:  
(A) LENGTH: 20 base pairs  
(B) TYPE: nucleic acid  
(C) STRANDEDNESS: single  
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

25 (iv) ANTI-SENSE: NO

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:17:

CCCACCTTTT GAGCAAGTTC

20

30 (2) INFORMATION FOR SEQ ID NO:18:

- (i) SEQUENCE CHARACTERISTICS:  
(A) LENGTH: 20 base pairs  
(B) TYPE: nucleic acid  
(C) STRANDEDNESS: single  
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

(iv) ANTI-SENSE: NO

5

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:18:

GGCTTATGAG TATTCTTCC

20

(2) INFORMATION FOR SEQ ID NO:19:

10

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 326 amino acids
- (B) TYPE: amino acid
- (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: protein

(iii) HYPOTHETICAL: NO

15

(iv) ANTI-SENSE: NO

(v) FRAGMENT TYPE: N-terminal

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:19:

20

Met Pro Pro Ser Ile Ser Ala Phe Gln Ala Ala Tyr Ile Gly Ile Glu  
1 5 10 15

Val Leu Ile Ala Leu Val Ser Val Pro Gly Asn Val Leu Val Ile Trp  
20 25 30

Ala Val Lys Val Asn Gln Ala Leu Arg Asp Ala Thr Phe Cys Phe Ile  
35 40 45

25

Val Ser Leu Ala Val Ala Asp Val Ala Val Gly Ala Leu Val Ile Pro  
50 55 60

Leu Ala Ile Leu Ile Asn Ile Gly Pro Gln Thr Tyr Phe His Thr Cys  
65 70 75 80

30

Leu Met Val Ala Cys Pro Val Leu Ile Leu Thr Gln Ser Ser Ile Leu  
85 90 95

Ala Leu Leu Ala Ile Ala Val Asp Arg Tyr Leu Arg Val Lys Ile Pro  
100 105 110

Leu Arg Tyr Lys Met Val Val Thr Pro Arg Arg Ala Ala Val Ala Ile  
115 120 125

Ala Gly Cys Trp Ile Leu Ser Phe Val Val Gly Leu Thr Pro Met Phe  
130 135 140

Gly Trp Asn Asn Leu Ser Ala Val Glu Arg Ala Trp Ala Ala Asn Gly  
145 150 155 160

Ser Met Gly Glu Pro Val Ile Lys Cys Glu Phe Glu Lys Val Ile Ser  
165 170 175

5 Met Glu Tyr Met Val Tyr Phe Asn Phe Phe Val Trp Val Leu Pro Pro  
180 185 190

Leu Leu Leu Met Val Leu Ile Tyr Leu Glu Val Phe Tyr Leu Ile Arg  
195 200 205

10 Lys Gln Leu Asn Lys Lys Val Ser Ala Ser Ser Gly Asp Pro Gln Lys  
210 215 220

Tyr Tyr Gly Lys Glu Leu Lys Ile Ala Lys Ser Leu Ala Leu Ile Leu  
225 230 235 240

Phe Leu Phe Ala Leu Ser Trp Leu Pro Leu His Ile Leu Asn Cys Ile  
245 250 255

15 Thr Leu Phe Cys Pro Ser Cys His Lys Pro Ser Ile Leu Thr Tyr Ile  
260 265 270

Ala Ile Phe Leu Thr His Gly Asn Ser Ala Met Asn Pro Ile Val Tyr  
275 280 285

Ala Phe Arg Ile Gln Lys Phe Arg Val Thr Phe Leu Lys Ile Trp Asn  
290 295 300

20 Asp His Phe Arg Cys Gln Pro Ala Pro Pro Ile Asp Glu Asp Leu Pro  
305 310 315 320

Glu Glu Arg Pro Asp Asp  
325

(2) INFORMATION FOR SEQ ID NO:20:

- 25 (i) SEQUENCE CHARACTERISTICS:  
(A) LENGTH: 981 base pairs  
(B) TYPE: nucleic acid  
(C) STRANDEDNESS: double  
(D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: cDNA
- 30 (iii) HYPOTHETICAL: NO
- (iv) ANTI-SENSE: NO

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:20:

|             |            |            |             |            |             |            |     |
|-------------|------------|------------|-------------|------------|-------------|------------|-----|
| ATGCCGCCCT  | CCATCTCAGC | TTTCCAGGCC | GCCTACATCG  | GCATCGAGGT | GCTCATCGCC  | 60         |     |
| CTGGTCTCTG  | TGCCCCGGAA | CGTGCTGGTG | ATCTGGCGG   | TGAAGGTGAA | CCAGGGCGCTG | 120        |     |
| CGGGATGCCA  | CCTTCTGCTT | CATCGTGTG  | CTGGCGGTGG  | CTGATGTGGC | CGTGGGTGCC  | 180        |     |
| 5           | CTGGTCATCC | CCCTCGCCAT | CCTCATCAAC  | ATTGGGCCAC | AGACCTACTT  | CCACACCTGC | 240 |
| CTCATGGTTG  | CCTGTCCGGT | CCTCATCCTC | ACCCAGAGCT  | CCATCCTGGC | CCTGCTGGCA  | 300        |     |
| ATIGCTGTGG  | ACCGCTACCT | CCGGGTCAAG | ATCCCTCTCC  | GGTACAAGAT | GGTGGTGACC  | 360        |     |
| CCCCGGAGGG  | CGGCGGTGGC | CATAGCCGGC | TGCTGGATCC  | TCTCCTTCGT | GGTGGGACTG  | 420        |     |
| 10          | ACCCCTATGT | TTGGCTGGAA | CAATCTGAGT  | GCGGTGGAGC | GGGCCTGGGC  | AGCCAACGGC | 480 |
| ACCATGGGGG  | AGCCCCTGAT | CAAATGCGAG | TTCGAGAAGG  | TCATCAGCAT | GGAGTACATG  | 540        |     |
| GTCTACTTCA  | ACTTCTTTGT | GTGGGTGCTG | CCCCCGCTTC  | TCCTCATGGT | CCTCATCTAC  | 600        |     |
| CTGGAGGTCT  | TCTACCTAAT | CCGCAAGCAG | CTCAACAAGA  | AGGTGTCCGC | CTCCTCCGGC  | 660        |     |
| 15          | GACCCGCAGA | AGTACTATGG | GAAGGGAGCTG | AAGATCGCCA | AGTCGCTGGC  | CCTCATCCTC | 720 |
| TTCCCTTTG   | CCCTCAGCTG | GCTGCCTTG  | CACATCCTCA  | ACTGCATCAC | CCTCTTCTGC  | 780        |     |
| CCGTCCCTGCC | ACAAGCCCAG | CATCCTTACC | TACATTGCCA  | TCTTCCTCAC | GCACGGCAAC  | 840        |     |
| TCGGCCATGA  | ACCCCATTGT | CTATGCCTTC | CGCATCCAGA  | AGTTCCGCGT | CACCTTCCTT  | 900        |     |
| AAGATTGGA   | ATGACCATTT | CCGCTGCCAG | CCTGCACCTC  | CCATTGACGA | GGATCTCCCA  | 960        |     |
| 20          | GAAGAGAGGC | CTGATGACTA | G           |            |             | 981        |     |

(2) INFORMATION FOR SEQ ID NO:21:

- (i) SEQUENCE CHARACTERISTICS:  
(A) LENGTH: 412 amino acids  
(B) TYPE: amino acid  
25 (D) TOPOLOGY: linear  
  
(ii) MOLECULE TYPE: protein  
  
(iii) HYPOTHETICAL: NO  
  
(iv) ANTI-SENSE: NO  
  
30 (v) FRAGMENT TYPE: N-terminal

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:21:

|     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |     |
|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|
| Met | Pro | Ile | Met | Gly | Ser | Ser | Val | Tyr | Ile | Thr | Val | Glu | Leu | Ala | Ile |
| 1   |     |     |     |     | 5   |     |     |     |     |     | 10  |     |     |     | 15  |

Ala Val Leu Ala Ile Leu Gly Asn Val Leu Val Cys Trp Ala Val Trp  
20 25 30

Leu Asn Ser Asn Leu Gln Asn Val Thr Asn Tyr Phe Val Val Ser Leu  
35 40 45

5 Ala Ala Ala Asp Ile Ala Val Gly Val Leu Ala Ile Pro Phe Ala Ile  
50 55 60

Thr Ile Ser Thr Gly Phe Cys Ala Ala Cys His Gly Cys Leu Phe Ile  
65 70 75 80

Ala Cys Phe Val Leu Val Leu Thr Gln Ser Ser Ile Phe Ser Leu Leu  
85 90 95

10 Ala Ile Ala Ile Asp Arg Tyr Ile Ala Ile Arg Ile Pro Leu Arg Tyr  
100 105 110

Asn Gly Leu Val Thr Gly Thr Arg Ala Lys Gly Ile Ile Ala Ile Cys  
115 120 125

Trp Val Leu Ser Phe Ala Ile Gly Leu Thr Pro Met Leu Gly Trp Asn  
130 135 140

15 Asn Cys Gly Gln Pro Lys Glu Gly Lys Asn His Ser Gln Gly Cys Gly  
145 150 155 160

Glu Gly Gln Val Ala Cys Leu Phe Glu Asp Val Val Pro Met Asn Tyr  
165 170 175

Met Val Tyr Phe Asn Phe Phe Ala Cys Val Leu Val Pro Leu Leu Leu  
180 185 190

20 Met Leu Gly Val Tyr Leu Arg Ile Phe Leu Ala Ala Arg Arg Gln Leu  
195 200 205

Lys Gln Met Glu Ser Gln Pro Leu Pro Gly Glu Arg Ala Arg Ser Thr  
210 215 220

25 Leu Gln Lys Glu Val His Ala Ala Lys Ser Leu Ala Ile Ile Val Gly  
225 230 235 240

Leu Phe Ala Leu Cys Trp Leu Pro Leu His Ile Ile Asn Cys Phe Thr  
245 250 255

Phe Phe Cys Pro Asp Cys Ser His Ala Pro Leu Trp Leu Met Tyr Leu  
260 265 270

30 Ala Ile Val Leu Ser His Thr Asn Ser Val Val Asn Pro Phe Ile Tyr  
275 280 285

Ala Tyr Arg Ile Arg Glu Phe Arg Gln Thr Phe Arg Lys Ile Ile Arg  
290 295 300

Ser His Val Leu Arg Gln Gln Glu Pro Phe Lys Ala Ala Gly Thr Ser

|    |   |     |     |     |
|----|---|-----|-----|-----|
|    | 305   | 310 | 315 | 320 |
|    | Ala Arg Val Leu Ala Ala His Gly Ser Asp Gly Glu Gln Val Ser Leu |     |     |     |
|    | 325   |     | 330 | 335 |
|    | Arg Leu Asn Gly His Pro Pro Gly Val Trp Ala Asn Gly Ser Ala Pro |     |     |     |
|    | 340   |     | 345 | 350 |
| 5  | His Pro Glu Arg Arg Pro Asn Gly Tyr Ala Leu Gly Leu Val Ser Gly |     |     |     |
|    | 355   |     | 360 | 365 |
|    | Gly Ser Ala Gln Glu Ser Gln Gly Asn Thr Gly Leu Pro Asp Val Glu |     |     |     |
|    | 370   |     | 375 | 380 |
|    | Leu Leu Ser His Glu Leu Lys Gly Val Cys Pro Glu Pro Pro Gly Leu |     |     |     |
|    | 385   |     | 390 | 395 |
| 10 | Asp Asp Pro Leu Ala Gln Asp Gly Ala Gly Val Ser                 |     |     |     |
|    | 405   |     | 410 |     |

(2) INFORMATION FOR SEQ ID NO:22:

- (i) SEQUENCE CHARACTERISTICS:
- (A) LENGTH: 1239 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: double
- (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: cDNA
- (iii) HYPOTHETICAL: NO
- (iv) ANTI-SENSE: NO

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:22:

|    |  |     |
|----|--|-----|
|    | ATGCCCATCA TGGGCTCCTC GGTGTACATC ACGGTGGAGC TGGCCATTGC TGTGCTGGCC  | 60  |
| 25 | ATCCTGGGCA ATGTGCTGGT GTGCTGGCC GTGTGGCTCA ACAGCAACCT GCAGAACGTC   | 120 |
|    | ACCAAACACT TTGTGGTGTC ACTGGCGGCG GCCGACATCG CAGTGGGTGT GCTCGCCATC  | 180 |
|    | CCCTTTGCCA TCACCATCAG CACCGGGTTC TGCGCTGCCT GCCACGGCTG CCTCTTCATT  | 240 |
|    | GCCTGCTTCG TCCTGGTCCT CACGCAGAGC TCCATCTTCA GTCTCCTGGC CATGCCATT   | 300 |
| 30 | GACCGCTACA TTGCCATCCG CATCCCCGTC CGGTACAATG GCTTGGTGAC CGGCACGAGG  | 360 |
|    | GCTAAGGGCA TCATTGCCAT CTGCTGGGTG CTGTCGTTTG CCATCGGCCT GACTCCCATG  | 420 |
|    | CTAGGTTGGA ACAACTGCGG TCAGCCAAG GAGGGCAAGA ACCACTCCCCA GGGCTGCGGG  | 480 |
|    | GAGGGCCAAG TGGCCTGTCT CTTTGAGGAT GTGGTCCCCA TGAACATACAT GGTGTACTTC | 540 |

|   |      |
|---|------|
| AACTTCTTG CCTGTGTGCT GGTGCCCTG CTGCTCATGC TGGGTGTCTA TTTGCGGATC     | 600  |
| TTCCTGGCGG CGCGACGACA GCTGAAGCAG ATGGAGAGCC AGCCTCTGCC GGGGGAGCGG   | 660  |
| GCACGGTCCA CACTGCAGAA GGAGGTCCAT GCTGCCAAGT CACTGCCAT CATTGTGGGG    | 720  |
| CTCTTGCCC TCTGCTGGCT GCCCCTACAC ATCATCAACT GCTTCACTTT CTTCTGCC      | 780  |
| 5 GACTGCAGCC ACGCCCCTCT CTGGCTCATG TACCTGGCCA TCGTCCTCTC CCACACCAAT | 840  |
| TCGGTTGTGA ATCCCTTCAT CTACGCCTAC CGTATCCGCG AGTTCCGCCA GACCTTCCGC   | 900  |
| AAGATCATTC GCAGCCACGT CCTGAGGCAG CAAGAACCTT TCAAGGCAGC TGGCACCAGT   | 960  |
| GCCCCGGTCT TGGCAGCTCA TGGCAGTGAC GGAGAGCAGG TCAGCCTCCG TCTCAACGGC   | 1020 |
| 10 CACCCGCCAG GAGTGTGGC CAACGGCA GTCTCCCCACC CTGAGCGGAG GCCCAATGGC  | 1080 |
| TATGCCCTGG GGCTGGTGAG TGGAGGGAGT GCCCAAGAGT CCCAGGGAA CACGGGCCTC    | 1140 |
| CCAGACGTGG AGCTCCTTAG CCATGAGCTC AAGGGAGTGT GCCCAGAGCC CCCTGGCCTA   | 1200 |
| GATGACCCCCC TGGCCCAGGA TGGAGCCAGGA GTGTCC                           | 1239 |

15 (2) INFORMATION FOR SEQ ID NO:23:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 332 amino acids
- (B) TYPE: amino acid
- (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: protein

20 (iii) HYPOTHETICAL: NO

(iv) ANTI-SENSE: NO

(v) FRAGMENT TYPE: N-terminal

25 (ix) FEATURE:

- (A) NAME/KEY: Modified-site
- (B) LOCATION: 216
- (D) OTHER INFORMATION: /label= threonine

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:23:

|  |    |    |    |
|--|----|----|----|
| 30 Met Leu Leu Glu Thr Gln Asp Ala Leu Tyr Val Ala Leu Glu Leu Val |    |    |    |
| 1  | 5  | 10 | 15 |
| Ile Ala Ala Leu Ser Val Ala Gly Asn Val Leu Val Cys Ala Ala Val    |    |    |    |
| 20   | 25 | 30 |    |
| Gly Thr Ala Asn Thr Leu Gln Thr Pro Thr Asn Tyr Phe Leu Val Ser    |    |    |    |
| 35   | 40 | 45 |    |

Leu Ala Ala Ala Asp Val Ala Val Gly Leu Phe Ala Ile Pro Phe Ala  
50 55 60

Ile Thr Ile Ser Leu Gly Phe Cys Thr Asp Phe Tyr Gly Cys Leu Phe  
65 70 75 80

5 Leu Ala Cys Phe Val Leu Val Leu Thr Gln Ser Ser Ile Phe Ser Leu  
85 90 95

Leu Ala Val Ala Val Asp Arg Tyr Leu Ala Ile Cys Val Pro Leu Arg  
100 105 110

Tyr Lys Ser Leu Val Thr Gly Thr Arg Ala Arg Gly Val Ile Ala Val  
115 120 125

10 Leu Trp Val Leu Ala Phe Gly Ile Gly Leu Thr Pro Phe Leu Gly Trp  
130 135 140

Asn Ser Lys Asp Ser Ala Thr Asn Asn Cys Thr Glu Pro Trp Asp Gly  
145 150 155 160

Thr Thr Asn Glu Ser Cys Cys Leu Val Lys Cys Leu Phe Glu Asn Val  
165 170 175

15 Val Pro Met Ser Tyr Met Val Tyr Phe Asn Phe Phe Gly Cys Val Leu  
180 185 190

Pro Pro Leu Leu Ile Met Leu Val Ile Tyr Ile Lys Ile Phe Leu Val  
195 200 205

Ala Cys Arg Gln Leu Gln Arg Xaa Glu Leu Met Asp His Ser Arg Thr  
210 215 220

20 Thr Leu Gln Arg Glu Ile His Ala Ala Lys Ser Leu Ala Met Ile Val  
225 230 235 240

Gly Ile Phe Ala Leu Cys Trp Leu Pro Val His Ala Val Asn Cys Val  
245 250 255

25 Thr Leu Phe Gln Pro Ala Gln Gly Lys Asn Lys Pro Lys Trp Ala Met  
260 265 270

Asn Met Ala Ile Leu Leu Ser His Ala Asn Ser Val Val Asn Pro Ile  
275 280 285

Val Tyr Ala Tyr Arg Asn Arg Asp Phe Arg Tyr Thr Phe His Lys Ile  
290 295 300

30 Ile Ser Arg Tyr Leu Leu Cys Gln Ala Asp Val Lys Ser Gly Asn Gly  
305 310 315 320

Gln Ala Gly Val Gln Pro Ala Leu Gly Val Gly Leu  
325 330

(2) INFORMATION FOR SEQ ID NO:24:

- (i) SEQUENCE CHARACTERISTICS:  
(A) LENGTH: 999 base pairs  
(B) TYPE: nucleic acid  
(C) STRANDEDNESS: double  
(D) TOPOLOGY: linear

- 5 (ii) MOLECULE TYPE: cDNA  
(iii) HYPOTHETICAL: NO  
(iv) ANTI-SENSE: NO

10 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:24:

|               |             |            |            |            |            |     |
|---------------|-------------|------------|------------|------------|------------|-----|
| ATGCTGCTGG    | AGACACAGGA  | CGCGCTGTAC | GTGGCGCTGG | AGCTGGTCAT | CGCCGCGCTT | 60  |
| TCGGTGGCGG    | GCAACGTGCT  | GGTGTGCGCC | GCGGTGGGCA | CGGGCAACAC | TCTGCAGACG | 120 |
| CCCACCAACT    | ACTTCCTGGT  | GTCCCTGGCT | GCGGCCGACG | TGGCCGTGGG | GCTCTTCGCC | 180 |
| ATCCCCCTTG    | CCATCACCAT  | CAGCCTGGGC | TTCTGCACTG | ACTTCTACGG | CTGCCTCTTC | 240 |
| 15 CTCGCCTGCT | TCGTGCTGGT  | GCTCACCGAG | AGCTCCATCT | TCAGCCTTCT | GGCCGTGGCA | 300 |
| GTCGACAGAT    | ACCTGGCCAT  | CTGTGTCCCG | CTCAGGTATA | AAAGTTGGT  | CACGGGGACC | 360 |
| CGAGCAAGAG    | GGGTCAATTGC | TGTCCTCTGG | GTCCTTGCCT | TTGGCATCGG | ATTGACTCCA | 420 |
| TTCCTGGGGT    | GGAACAGTAA  | AGACAGTGCC | ACCAACAACT | GCACAGAAC  | CTGGGATGGA | 480 |
| 20 ACCACGAATG | AAAGCTGCTG  | CCTTGTGAAG | TGTCTCTTTG | AGAATGTGGT | CCCCATGAGC | 540 |
| TACATGGTAT    | ATTTCAATT   | CTTTGGGTGT | GTTCTGCC   | CACTGCTTAT | AATGCTGGTG | 600 |
| ATCTACATTA    | AGATCTTCCT  | GGTGGCCTGC | AGGCAGCTTC | AGCGCACTGA | GCTGATGGAC | 660 |
| CACTCGAGGA    | CCACCCCTCCA | GCGGGAGATC | CATGCAGCCA | AGTCACTGGC | CATGATTGTG | 720 |
| 25 GGGATTTTG  | CCCTGTGCTG  | GTTACCTGTG | CATGCTGTTA | ACTGTGTCAC | TCTTTCCAG  | 780 |
| CCAGCTCAGG    | GTAAAAATAA  | GCCCAAGTGG | GCAATGAATA | TGGCCATTCT | TCTGTCACAT | 840 |
| GCCAATTCA     | TTGTCATCC   | CATTGTCTAT | GCTTACCGGA | ACCGAGACTT | CCGCTACACT | 900 |
| TTTCACAAA     | TTATCTCCAG  | GTATCTCTC  | TGCCAAGCAG | ATGTCAAGAG | TGGGAATGGT | 960 |
| 30 CAGGCTGGGG | TACAGCCTGC  | TCTCGGTGTG | GGCCTATGA  |            |            | 999 |

(2) INFORMATION FOR SEQ ID NO:25:

- (i) SEQUENCE CHARACTERISTICS:  
(A) LENGTH: 318 amino acids  
(B) TYPE: amino acid

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: protein

(iii) HYPOTHETICAL: NO

(iv) ANTI-SENSE: NO

5 (v) FRAGMENT TYPE: N-terminal

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:25:

Met Pro Asn Asn Ser Thr Ala Leu Ser Leu Ala Asn Val Thr Tyr Ile  
1 5 10 15

10 Thr Met Glu Ile Phe Ile Gly Leu Cys Ala Ile Val Gly Asn Val Leu  
20 25 30

Val Ile Cys Val Val Lys Leu Asn Pro Ser Leu Gln Thr Thr Phe  
35 40 45

15 Tyr Phe Ile Val Ser Leu Ala Leu Ala Asp Ile Ala Val Gly Val Leu  
50 55 60

Val Met Pro Leu Ala Ile Val Val Ser Leu Gly Ile Thr Ile His Phe  
65 70 75 80

Tyr Ser Cys Leu Phe Met Thr Cys Leu Leu Ile Phe Thr His Ala  
85 90 95

20 Ser Ile Met Ser Leu Leu Ala Ile Ala Val Asp Arg Tyr Leu Arg Val  
100 105 110

Lys Leu Thr Val Arg Tyr Lys Arg Val Thr Thr His Arg Arg Ile Trp  
115 120 125

Leu Ala Leu Gly Leu Cys Trp Leu Val Ser Phe Leu Val Gly Leu Thr  
130 135 140

25 Pro Met Phe Gly Trp Asn Met Lys Leu Thr Ser Glu Tyr His Arg Asn  
145 150 155 160

Val Thr Phe Leu Ser Cys Gln Phe Val Ser Val Met Arg Met Asp Tyr  
165 170 175

Met Val Tyr Phe Ser Phe Leu Thr Trp Ile Phe Ile Pro Leu Val Val  
180 185 190

30 Met Cys Ala Ile Tyr Leu Asp Ile Phe Tyr Ile Ile Arg Asn Lys Leu  
195 200 205

Ser Leu Asn Leu Ser Asn Ser Lys Glu Thr Gly Ala Phe Tyr Gly Arg  
210 215 220

Glu Phe Lys Thr Ala Lys Ser Leu Phe Leu Val Leu Phe Leu Phe Ala  
225 230 235 240

Leu Ser Trp Leu Pro Leu Ser Ile Ile Asn Cys Ile Ile Tyr Phe Asn  
245 250 255

Gly Glu Val Pro Gln Leu Val Leu Tyr Met Gly Ile Leu Leu Ser His  
260 265 270

5 Ala Asn Ser Met Met Asn Pro Ile Val Tyr Ala Tyr Lys Ile Lys Lys  
275 280 285

Phe Lys Glu Thr Tyr Leu Leu Ile Leu Lys Ala Cys Val Val Cys His  
290 295 300

10 Pro Ser Asp Ser Leu Asp Thr Ser Ile Glu Lys Asn Ser Glu  
305 310 315

(2) INFORMATION FOR SEQ ID NO:26:

- (i) SEQUENCE CHARACTERISTICS:  
(A) LENGTH: 957 base pairs  
(B) TYPE: nucleic acid  
(C) STRANDEDNESS: double  
15 (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

(iv) ANTI-SENSE: NO

20

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:26:

ATGCCCAACA ACAGCACTGC TCTGTCATTG GCCAATGTTA CCTACATCAC CATGGAAATT 60  
TTCATTGGAC TCTGCGCCAT AGTGGGCAAC GTGCTGGTCA TCTGCGTGGT CAAGCTGAAC 120  
25 CCCAGCCTGC AGACCACCCAC CTTCTATTTTC ATTCCTCTC TAGCCCTGGC TGACATTGCT 180  
GTTGGGGTGC TGGTCATGCC TTTGGCCATT GTTGTCAAGCC TGGGCATCAC AATCCACTTC 240  
TACAGCTGCC TTTTTATGAC TTGCCTACTG CTTATCTTTA CCCACGCCCTC CATCATGTCC 300  
TTGCTGGCCA TCGCTGTGGA CCGATACTTG CGGGTCAAGC TTACCGTCAG ATACAAGAGG 360  
30 GTCACCACTC ACAGAAGAAT ATGGCTGGCC CTGGGCCTTT GCTGGCTGGT GTCATTCCCTG 420  
GTGGGATTGA CCCCCATGTT TGGCTGGAAC ATGAAACTGA CCTCAGAGTA CCACAGAAAT 480  
GTCACCTTCC TTTCATGCCA ATTTGTTCC GTCATGAGAA TGGACTACAT GGTATACTTC 540  
AGCTTCCTCA CCTGGATTTT CATCCCCCTG GTTGTCAATGT GCGCCATCTA TCTTGACATC 600

|            |            |            |            |            |            |            |     |
|------------|------------|------------|------------|------------|------------|------------|-----|
| TTTTACATCA | TTCGGAACAA | ACTCAGTCTG | AACTTATCTA | ACTCCAAAGA | GACAGGTGCA | 660        |     |
| TTTTATGGAC | GGGAGTTCAA | GACGGCTAAC | TCCTTGTTC  | TGGTTCTTTT | CTTGTGCT   | 720        |     |
| CTGTCATGGC | TGCCTTATC  | TATCATCAAC | TGCATCATCT | ACTTTAATGG | TGAGGTACCA | 780        |     |
| CAGCTTGTGC | TGTACATGGG | CATCCTGCTG | TCCCAGGCCA | ACTCCATGAT | GAACCCTATC | 840        |     |
| 5          | GTCTATGCCT | ATAAAATAAA | GAAGTTCAAG | GAAACCTACC | TTTTGATCCT | CAAAGCCTGT | 900 |
|            | GTGGTCTGCC | ATCCCTCTGA | TTCTTGAC   | ACAAGCATTG | AGAAGAATTG | TGAGTAG    | 957 |

(2) INFORMATION FOR SEQ ID NO:27:

- 10 (i) SEQUENCE CHARACTERISTICS:  
(A) LENGTH: 27 base pairs  
(B) TYPE: nucleic acid  
(C) STRANDEDNESS: single  
(D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: cDNA
- (iii) HYPOTHETICAL: NO
- 15 (iv) ANTI-SENSE: NO

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:27:

CCCAAGCTTA TGAAAGCCAA CAATACC

27

20 (2) INFORMATION FOR SEQ ID NO:28:

- (i) SEQUENCE CHARACTERISTICS:  
(A) LENGTH: 27 base pairs  
(B) TYPE: nucleic acid  
(C) STRANDEDNESS: single  
(D) TOPOLOGY: linear

25 (ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

(iv) ANTI-SENSE: NO

30 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:28:

TGCTCTAGAC TCTGGTATCT TCACATT

27

(2) INFORMATION FOR SEQ ID NO:29:

- (i) SEQUENCE CHARACTERISTICS:  
(A) LENGTH: 45 base pairs

- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: both
- (D) TOPOLOGY: linear

5 (ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

5 (iv) ANTI-SENSE: YES

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:29:

GCCTCTTTGA GGATGTGGTC CCCATGAAC ACATGGTGTA CTTCA

45

10 (2) INFORMATION FOR SEQ ID NO:30:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 45 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: both
  - (D) TOPOLOGY: linear

15 (ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

(iv) ANTI-SENSE: YES

20 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:30:

GCAGGGGCAC CAGCACACAG GCAAAGAAAGT TGAAGTACAC CATGT

45

(2) INFORMATION FOR SEQ ID NO:31:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 18 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: both
  - (D) TOPOLOGY: linear

25 (ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

30 (iv) ANTI-SENSE: YES

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:31:

TCACCACATT CCAGGAGC

18

(2) INFORMATION FOR SEQ ID NO:32:

- (i) SEQUENCE CHARACTERISTICS:  
(A) LENGTH: 18 base pairs  
(B) TYPE: nucleic acid  
(C) STRANDEDNESS: both  
(D) TOPOLOGY: linear

5 (ii) MOLECULE TYPE: cDNA  
(iii) HYPOTHETICAL: NO  
(iv) ANTI-SENSE: YES

10 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:32:

ACTCCCTTGGA GGCCATGT

18

(2) INFORMATION FOR SEQ ID NO:33:

- (i) SEQUENCE CHARACTERISTICS:  
(A) LENGTH: 45 base pairs  
15 (B) TYPE: nucleic acid  
(C) STRANDEDNESS: both  
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA  
(iii) HYPOTHETICAL: NO  
20 (iv) ANTI-SENSE: YES

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:33:

TCCTGCACCA CCAAAGTCTT AGCCCCCTG GCCAAGGTCA TCCAT

45

25 (2) INFORMATION FOR SEQ ID NO:34:

- (i) SEQUENCE CHARACTERISTICS:  
(A) LENGTH: 45 base pairs  
(B) TYPE: nucleic acid  
(C) STRANDEDNESS: both  
(D) TOPOLOGY: linear

30 (ii) MOLECULE TYPE: cDNA  
(iii) HYPOTHETICAL: NO  
(iv) ANTI-SENSE: YES

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:34:

CATGAGCCCT TCCACGATGC CAAAGTTGTC ATGGATGACC TTGGC

45

(2) INFORMATION FOR SEQ ID NO:35:

(i) SEQUENCE CHARACTERISTICS:

- 5 (A) LENGTH: 18 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: both
- (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

10 (iv) ANTI-SENSE: YES

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:35:

GTTACCTACA TCACCATG

18

15 (2) INFORMATION FOR SEQ ID NO:36:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 18 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: both
- (D) TOPOLOGY: linear

20 (ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

(iv) ANTI-SENSE: YES

25 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:36:

GTTAGATAAG TTCAGACT

18

(2) INFORMATION FOR SEQ ID NO:37:

(i) SEQUENCE CHARACTERISTICS:

- 30 (A) LENGTH: 45 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: both
- (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

(iv) ANTI-SENSE: YES

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:37:

CTGACCTCAG AGTACCACAG AAATGTCACC TTCCTTCAT GCCAA

45

5 (2) INFORMATION FOR SEQ ID NO:38:

- (i) SEQUENCE CHARACTERISTICS:  
(A) LENGTH: 45 base pairs  
(B) TYPE: nucleic acid  
(C) STRANDEDNESS: both  
(D) TOPOLOGY: linear

10 (ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

(iv) ANTI-SENSE: YES

15 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:38:

TTGGCATGAA AGGAAGGTGA CATTCTGTG GTACTCTGAG GTCAG

45

(2) INFORMATION FOR SEQ ID NO:39:

- (i) SEQUENCE CHARACTERISTICS:  
(A) LENGTH: 46 base pairs  
20 (B) TYPE: nucleic acid  
(C) STRANDEDNESS: both  
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

25 (iv) ANTI-SENSE: YES

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:39:

CTCAGTCTGA ACTTATCTAA CTCCAAAGAG ACAGGTGCAT TTTATG

46

30 (2) INFORMATION FOR SEQ ID NO:40:

- (i) SEQUENCE CHARACTERISTICS:  
(A) LENGTH: 46 base pairs  
(B) TYPE: nucleic acid  
(C) STRANDEDNESS: both  
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

(iv) ANTI-SENSE: YES

5 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:40:

CATAAAATGC ACCTGTCTCT TTGGAGTTAG ATAAGTTCACTGAG

46

(2) INFORMATION FOR SEQ ID NO:41:

(i) SEQUENCE CHARACTERISTICS:

- 10 (A) LENGTH: 18 base pairs  
(B) TYPE: nucleic acid  
(C) STRANDEDNESS: both  
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

15 (iv) ANTI-SENSE: YES

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:41:

TCCCTCGGTGT ACATCACG

18

20 (2) INFORMATION FOR SEQ ID NO:42:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 18 base pairs  
(B) TYPE: nucleic acid  
(C) STRANDEDNESS: both  
(D) TOPOLOGY: linear

25 (ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

(iv) ANTI-SENSE: YES

30 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:42:

TCCATCTGCT TCAGCTGT

18

(2) INFORMATION FOR SEQ ID NO:43:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 45 base pairs

- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: both
- (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

5 (iv) ANTI-SENSE: YES

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:43:

CTGGGCCTTT GCTGGCTGGT GTCATTCTTG GTGGGATTGA CCCCC

45

10 (2) INFORMATION FOR SEQ ID NO:44:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 45 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: both
  - (D) TOPOLOGY: linear

15 (ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

(iv) ANTI-SENSE: YES

20 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:44:

TGAGGTCAGT TTCATGTTCC AGCCAAACAT GGGGGTCAAT CCCAC

45

(2) INFORMATION FOR SEQ ID NO:45:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 20 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: both
  - (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

25 (iv) ANTI-SENSE: YES

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:45:

ATGCTGCTGG AGACACAGGA

20

(2) INFORMATION FOR SEQ ID NO:46:

- (i) SEQUENCE CHARACTERISTICS:  
(A) LENGTH: 19 base pairs  
(B) TYPE: nucleic acid  
(C) STRANDEDNESS: both  
(D) TOPOLOGY: linear

5 (ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

(iv) ANTI-SENSE: YES

10 (xi) SEQUENCE DESCRIPTION: SEQ ID NO:46:

TGGTCCATCA GCTCAGTGC

19

(2) INFORMATION FOR SEQ ID NO:47:

- (i) SEQUENCE CHARACTERISTICS:  
(A) LENGTH: 60 base pairs  
15 (B) TYPE: nucleic acid  
(C) STRANDEDNESS: both  
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

20 (iv) ANTI-SENSE: YES

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:47:

GGTGGAACAG TAAAGACAGT GCCACCAACA ACTGCACAGA ACCCTGGGAT GGAACCACGA

60

25 (2) INFORMATION FOR SEQ ID NO:48:

- (i) SEQUENCE CHARACTERISTICS:  
(A) LENGTH: 60 base pairs  
(B) TYPE: nucleic acid  
(C) STRANDEDNESS: both  
(D) TOPOLOGY: linear

30 (ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

(iv) ANTI-SENSE: YES

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:48:

GGACCACATT CTCAAAGAGA CACTTCACAA GGCAGCAGCT TTCATTCTGTG GTTCCATCCC

60

(2) INFORMATION FOR SEQ ID NO:49:

5

- (i) SEQUENCE CHARACTERISTICS:  
(A) LENGTH: 18 base pairs  
(B) TYPE: nucleic acid  
(C) STRANDEDNESS: both  
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

10

- (iii) HYPOTHETICAL: NO  
(iv) ANTI-SENSE: YES

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:49:

15

CTACATCGGC ATCGAGGT

18

(2) INFORMATION FOR SEQ ID NO:50:

20

- (i) SEQUENCE CHARACTERISTICS:  
(A) LENGTH: 18 base pairs  
(B) TYPE: nucleic acid  
(C) STRANDEDNESS: both  
(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

(iv) ANTI-SENSE: YES

25

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:50:

GAACTCGCAC TTGATCAC

18

(2) INFORMATION FOR SEQ ID NO:51:

30

- (i) SEQUENCE CHARACTERISTICS:  
(A) LENGTH: 45 base pairs  
(B) TYPE: nucleic acid  
(C) STRANDEDNESS: both  
(D) TOPOLOGY: both

(ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

(iv) ANTI-SENSE: YES

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:51:

5 TGGTGGGACT GACCCCTATG TTTGGCTGGA ACAATCTGAG TGCGG

45

(2) INFORMATION FOR SEQ ID NO:52:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 45 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: both
- (D) TOPOLOGY: linear

10

(ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

(iv) ANTI-SENSE: YES

15

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:52:

TGCTGCCGTT GGCTGCCAG GCCCGCTCCA CCGCACTCAG ATTGT

45

(2) INFORMATION FOR SEQ ID NO:53:

20

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 42 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: both
- (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

25

(iii) HYPOTHETICAL: NO

(iv) ANTI-SENSE: YES

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:53:

30

CTGAGCTCAG CAGACGAAAA CCTCACCTTC CTACCCTGCC GA

42

(2) INFORMATION FOR SEQ ID NO:54:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 42 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: both

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

(iv) ANTI-SENSE: YES

5

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:54:

TCGGCAGGGT AGGAAGGTGA GGTTTTCTGC TGCTGAGCTC AG

42

(2) INFORMATION FOR SEQ ID NO:55:

10

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 46 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: both
- (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

15

(iii) HYPOTHETICAL: NO

(iv) ANTI-SENSE: YES

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:55:

20

CTCAGCCAGA GCTTTCTGG CTCCAGAGAG ACAGGCGCAT TCTATG

46

(2) INFORMATION FOR SEQ ID NO:56:

25

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 46 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: both
- (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(iii) HYPOTHETICAL: NO

(iv) ANTI-SENSE: YES

30

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:56:

CATAGAACATGC GCCTGTCTCT CTGGAGCCAG AAAAGCTCTG GCTGAG

46

WHAT IS CLAIMED IS:

- 5        1. A method for inhibiting TNF $\alpha$  production which comprises contacting the A2b subtype of the adenosine receptor with an adenosine receptor agonist.
- 10      2. A method for treating or preventing autoimmune diseases including rheumatoid arthritis, rheumatoid spondylitis, inflammatory bowel disease (ulcerative colitis and Crohns disease), intestinal pathology associated with graft vs. host disease, organ transplant reactions, septic shock, fever and myalgia due to infection and cachexia associated with chronic infections, malignancy and acquired immune deficiency syndrome, pulmonary diseases such as pulmonary sarcoidosis, silicosis, chronic pulmonary inflammatory disease, adult respiratory distress syndrome which comprises providing a sufficient quantity of an A2b adenosine receptor agonist to inhibit TNF $\alpha$  production.
- 15      3. A method for increasing cAMP accumulation in monocytes, and thereby inhibiting production of TNF $\alpha$ , which comprises contacting the monocyte A2b adenosine receptor subtype with an adenosine receptor agonist at a sufficient concentration to activate adenylate cyclase.
- 20      4. The method of any one of claims 1, 2, 3, or 4, wherein the adenosine receptor agonist is adenosine, CPCA, NECA, R-PIA, or CHA.
- 25      5. A method for inhibiting TNF $\alpha$  production which comprises contacting the A2b subtype of the adenosine receptor with an A2b adenosine receptor enhancer.

6. A method for identifying A2b adenosine receptor agonist enhancer or A2b receptor selective compounds which comprises the steps of:

- (a) contacting monocytes with a test compound and measuring the effect of the test compound on TNF $\alpha$  production;
- 5 (b) contacting a test compound, identified according to step (a) as inhibiting TNF $\alpha$  production by the monocytes, with membranes derived from a stable cell line individually expressing each of the A1, A2a, A2b, or A3 adenosine receptor or with the whole cell expressing each of the A1, A2a, A2b, or A3 adenosine receptor and measuring the binding affinity of the test compound for the receptor or the effect of the test compound on cAMP production in the stable cell line;
- 10 (c) selecting compounds which bind to the A2b adenosine receptor or which induces elevation in cAMP in the cell line expressing the A2b adenosine receptor and which do not bind to membranes or affect the cAMP level in the stable cell lines expressing the A1, A2a, or A3 adenosine receptor subtypes.

7. A method for inhibiting production of TNF $\alpha$  by activated monocytes which comprises contacting monocytes with an inhibitorily effective amount of a compound identified according to  
20 Claim 6.

**Patents Act 1977****Examiner's report to the Comptroller under Section 17  
(The Search report)**

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| <b>Relevant Technical Fields</b><br>(i) UK Cl (Ed.N) A5B (BHA)<br>(ii) Int Cl (Ed.6) A61K 31/52 | <b>Application number</b><br><b>GB 9508844.9</b> |
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| <b>Search Examiner</b><br><b>C SHERRINGTON</b> |
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| <b>Date of completion of Search</b><br><b>16 AUGUST 1995</b> |
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| <b>Documents considered relevant</b><br><b>following a search in respect of</b><br><b>Claims :-</b><br><b>1, 3 to 7</b> |
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**Databases (see below)**

(i) UK Patent Office collections of GB, EP, WO and US patent specifications.

(ii) ONLINE: WPI, CLAIMS, DIALOG/BIOTECH

**Categories of documents**

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|----------|--|--|----------------------|
| X        | GB 2264948 A                                       | (MERCK & CO INC) whole document, especially Table L, page 8, line 25; page 11, line 7 to page 13, line 13    | 1, 3 to 7            |
| X        | WO 93/25677 A1                                     | (GARVAN INSTITUTE OF MEDICAL RESEARCH) whole document, especially page 3, lines 17 to 31, Claim 7; Figure 4B | 1, 3 to 7            |
| X        | US Pat. Appl. NTIS US 7-577528                     | especially pages 40 to 64; Figure 1  | 1, 3 to 5            |
| A        | Life Sci. 1993, 52, 1917-1924                      | Inhibition of human monocyte TNF production by adenosine receptor agonists                                   | 1, 4, 5              |
| X        | Biochem. Biophys. Res. Commun. 1992, 187(1), 86-93 | Molecular Cloning and Expression of an Adenosine A2b Receptor from Human Brain                               | 1, 3 to 7            |
| A        | Mol. Endocrinol. 1992, 6, 384-393                  | Molecular Cloning and Expression of the cDNA for a Novel A2-Adenosine Receptor Subtype                       | 1                    |

**Databases:** The UK Patent Office database comprises classified collections of GB, EP, WO and US patent specifications as outlined periodically in the Official Journal (Patents). The on-line databases considered for search are also listed periodically in the Official Journal (Patents).